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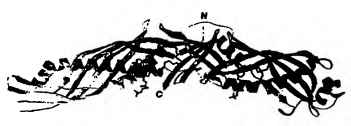
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(57) Abstract

The present invention solves the three-dimensional structure of BPI and thereby provides atomic coordinates of BPI from the analysis of x-ray diffraction patterns of sufficiently high resolution for three-dimensional structure determination of the protein, as well as methods for rational drug design, based on using amino acid sequence data and/or x-ray diffraction data provided on computer readable media, as analyzed on a computer system having suitable computer algorithms; and atomic coordinates are provided yielding structural information on related proteins, including the lipid binding and lipid transport protein family that includes BPI, LBP, CETP and PLTP.

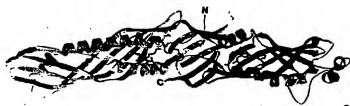


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BACTERICIDAL/PERMEABILITY-INCREASING PROTEIN: CRYSTALLIZATION, X-RAY DIFFRACTION, THREEDIMENSIONAL STRUCTURE DETERMINATION, RATIONAL DRUG DESIGN AND MOLECULAR MODELING OF RELATED PROTEINS

This is a continuation-in-part application of U.S. Application Serial No. 08/879,565, filed June 20, 1997, which is hereby incorporated by reference in its entirety.

The present invention generally pertains to the fields of protein crystallization, x-ray diffraction analysis, three-dimensional structural determination, rational drug design and molecular modeling of related proteins. The present invention solves the three-dimensional structure of bactericidal/permeability-increasing protein (BPI) and provides crystallization methods for BPI protein products. A crystallized BPI protein product was physically analyzed by x-ray diffraction techniques. The resulting x-ray diffraction patterns were of sufficiently high resolution to be useful for determining the three-dimensional structure of BPI and have yielded atomic coordinates for BPI. The present invention relates to uses of BPI coordinates for molecular modeling of related proteins and rational drug design (RDD) of mimetics and ligands for BPI and for related proteins. The present invention also relates to atomic coordinates of BPI, or portions thereof, to solve crystal forms of BPI proteins or their fragments, analogs, and variants thereof, or of related proteins, including lipid transfer proteins, or their fragments, analogs and variants.

Part of the work performed during development of this invention utilized U.S. Government funds. The U.S. Government has certain rights in this invention.

Background of the Invention

Bactericidal/permeability-increasing protein (BPI) is a protein isolated from the granules of mammalian polymorphonuclear leukocytes (PMNs or neutrophils), which are blood cells essential in the defense against invading microorganisms. BPI is known to bind the lipopolysaccharide (LPS), a major component of the outer membrane of gram-negative bacteria that stimulates a potent inflammatory response. Human BPI protein has been isolated from PMNs by acid extraction combined with either ion exchange chromatography [Elsbach, J. Biol. Chem., 254:11000 (1979)] or E. coli affinity chromatography [Weiss, et al., Blood, 69:652 (1987)]. BPI obtained

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in such a manner is referred to herein as natural BPI and has been shown to have potent bactericidal activity against a broad spectrum of gram-negative bacteria. The molecular weight of human BPI is approximately 55,000 daltons (55 kD). The amino acid sequence of the entire human BPI protein and the nucleic acid sequence of DNA encoding the protein have been reported in Figure 1 of Gray et al., *J. Biol. Chem.*, 264:9505 (1989), incorporated herein by reference. The Gray et al. amino acid sequence is set out in SEQ ID NO: 1 hereto. U.S. Patent No. 5,198,541, EP0375724 and WO89/10486 (PCT/US88/02700) disclose recombinant genes encoding and methods for expression of BPI proteins, including BPI holoprotein and fragments of BPI.

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A proteolytic N-terminal fragment of BPI having a molecular weight of about 25 kD possesses essentially all the anti-bacterial efficacy of the naturally-derived 55 kD human BPI holoprotein. [Ooi et al., *J. Bio. Chem.*, 262: 14891-14894 (1987)]. In contrast to the N-terminal portion, the C-terminal region of the isolated human BPI protein displays only slightly detectable anti-bacterial activity against gram-negative organisms. [Ooi et al., *J. Exp. Med.*, 174:649 (1991).] An N-terminal BPI fragment of approximately 23 kD, referred to as "rBPI₂₃," has been produced by recombinant means and also retains anti-bacterial activity against gram-negative organisms. [Gazzano-Santoro et al., *Infect. Immun.* 60:4754-4761 (1992).] An N-terminal analog of BPI, rBPI₂₁, has been produced as described in Horwitz et al., *Protein Expression Purification*, 8:28-40 (1996).

The bactericidal effect of BPI has been reported to be highly specific to gram-negative species, e.g., in Elsbach and Weiss, *Inflammation: Basic Principles and Clinical Correlates*, eds. Gallin et al., Chapter 30, Raven Press, Ltd. (1992). This reported target cell specificity was believed to be the result of the strong attraction of BPI for LPS on the outer membrane (or envelope) of gram-negative organisms. Although BPI was commonly thought to be non-toxic for other microorganisms, including yeast and for higher eukaryotic cells, it has recently been discovered that BPI protein products exhibit activity against gram-positive bacteria, mycoplasma, mycobacteria, fungi, protozoa and chlamydia.

The precise mechanism by which BPI kills gram-negative bacteria is not yet completely elucidated, but it is believed that BPI must first bind to the surface of the

bacteria through electrostatic and hydrophobic interactions between the cationic BPI protein and negatively charged sites on lipopolysaccharide. Bacterial LPS has been referred to as "endotoxin" because of the potent inflammatory response that it stimulates, i.e., the release of mediators by host inflammatory cells which may ultimately result in irreversible endotoxic shock. BPI binds to lipid A, reported to be the most toxic and most biologically active component of LPS.

In susceptible gram-negative bacteria, BPI binding is thought to disrupt LPS structure, leading to activation of bacterial enzymes that degrade phospholipids and peptidoglycans, altering the permeability of the cell's outer membrane, and initiating events that ultimately lead to cell death. [Elsbach and Weiss (1992), supra]. BPI has been proposed to act in two stages. The first stage proposed is a sublethal one that is characterized by immediate growth arrest, permeabilization of the outer membrane and selective activation of bacterial enzymes that hydrolyze phospholipids and peptidoglycans. Bacteria at this stage could be rescued by growth in serum albumin supplemented media [Mannion et al., J. Clin. Invest., 85:853-860 (1990)]. The second stage, defined by growth inhibition that cannot be reversed by serum albumin, is proposed to occur after prolonged exposure of the bacteria to BPI and characterized by extensive physiologic and structural changes, including apparent damage to the inner cytoplasmic membrane.

Initial binding of BPI to LPS leads to organizational changes that probably result from binding to the anionic groups of LPS, which normally stabilize the outer membrane through binding of Mg⁺⁺ and Ca⁺⁺. Attachment of BPI to the outer membrane of gram-negative bacteria produces rapid permeabilization of the outer membrane to hydrophobic agents such as actinomycin D. Binding of BPI and subsequent gram-negative bacterial killing depends, at least in part, upon the LPS polysaccharide chain length, with long O-chain bearing, "smooth" organisms being more resistant to BPI bactericidal effects than short O-chain bearing, "rough" organisms [Weiss et al., J. Clin. Invest. 65: 619-628 (1980)]. This permeabilization of the gram-negative outer envelope is reversible upon dissociation of the BPI, a process requiring high concentrations of divalent cations and synthesis of new LPS [Weiss et al., J. Immunol. 132: 3109-3115 (1984)]. Loss of gram-negative bacterial viability, however, is not reversed by processes which restore the envelope integrity,

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suggesting that the bactericidal action is mediated by additional lesions induced in the target organism and which may be situated at the cytoplasmic membrane (Mannion et al., J. Clin. Invest. 86: 631-641 (1990)). Specific investigation of this possibility has shown that on a molar basis BPI is at least as inhibitory of cytoplasmic membrane vesicle function as polymyxin B but the exact mechanism as well as the relevance of such vesicles to studies of intact organisms was not elucidated (In't Veld, et al., Infection and Immunity 56: 1203-1208 (1988)).

BPI is a member of a gene/protein family of lipopolysaccharide binding and lipid transfer proteins whose other currently known members include lipopolysaccharide binding protein (LBP), cholesteryl ester transfer protein (CETP) and phospholipid transfer protein (PLTP).

BPI protein products (which include naturally and recombinantly produced BPI protein; natural, synthetic, and recombinant biologically active polypeptide fragments of BPI protein; biologically active polypeptide variants of BPI protein or fragments thereof, including hybrid fusion proteins and dimers; biologically active polypeptide analogs of BPI protein or fragments or variants thereof, including cysteine-substituted analogs; and BPI-derived peptides) have been demonstrated to have a variety of beneficial activities. BPI protein products are known to be bactericidal for gramnegative bacteria, as described in U.S. Patent Nos. 5,198,541 and 5,523,288, both of which are incorporated herein by reference. BPI protein products are also known to enhance the effectiveness of antibiotic therapy in gram-negative bacterial infections, as described in U.S. Patent No. 5,523,288 and corresponding International Publication No. WO 95/08344 (PCT/US94/11225), which are incorporated herein by reference. BPI protein products are also known to be bactericidal for gram-positive bacteria and mycoplasma, and to enhance the effectiveness of antibiotics in grampositive bacterial infections, as described in U.S. Patent No. 5,578,572 and corresponding International Publication No. WO 95/19180 (PCT/US95/00656), which are incorporated herein by reference. BPI protein products are further known to exhibit anti-fungal activity, and to enhance the activity of other anti-fungal agents, as described in U.S. Patent No. 5,627,153 and corresponding International Publication No. WO 95/19179 (PCT/US95/00498), and further as described for anti-fungal peptides in co-pending U.S. Application Serial No. 08/621,259 filed March 21, 1996,

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which is in turn a continuation-in-part of U.S. Application Serial No. 08/504,841 filed July 20, 1994 and corresponding International Publication Nos. WO 96/08509 (PCT/US95/09262) and WO 97/04008 (PCT/US96/03845), all of which are incorporated herein by reference. BPI protein products are further known to exhibit anti-protozoan activity, as described in U.S. Patent No. 5,646,114 and corresponding International Publication No. WO 96/01647 (PCT/US95/08624), all of which are incorporated herein by reference. BPI protein products are known to exhibit antichlamydial activity, as described in co-pending U.S. Application Serial No. 08/694,843 filed August 9, 1996 and corresponding International Publication No. WO 98/06415 (PCT/US97/13810), all of which are incorporated herein by reference. Finally, BPI protein products are known to exhibit anti-mycobacterial activity, as described in co-pending U.S. Application Serial No. 08/626,646 filed April 1, 1996, which is in turn a continuation of U.S. Application Serial No. 08/285,803 filed August 14, 1994, which is in turn a continuation-in-part of U.S. Application Serial No. 08/031,145 filed March 12, 1993 and corresponding International Publication No. WO94/20129 (PCT/US94/02463), all of which are incorporated herein by reference.

The effects of BPI protein products in humans with endotoxin in circulation, including effects on TNF, IL-6 and endotoxin are described in U.S. Patent Nos. 5,643,875 and 5,753,620 and corresponding International Publication No. WO 95/19784 (PCT/US95/01151), all of which are incorporated herein by reference.

BPI protein products are also known to be useful for treatment of specific disease conditions, such as meningococcemia in humans (as described in co-pending U.S. Application Serial No. 08/644,287 filed May 10, 1996 and corresponding International Publication No. WO 97/42966 (PCT/US97/08016), which are incorporated herein by reference), hemorrhagic trauma in humans, (as described in co-pending U.S. Application Serial No. 08/862,785, a continuation-in-part of U.S. Serial No. 08/652,292 filed May 23, 1996, now U.S. Patent No. 5,756,464, and corresponding International Publication No. WO 97/44056 (PCT/US97/08941), all of which are incorporated herein by reference), burn injury (as described in U.S. Patent No. 5,494,896 and corresponding International Publication No. WO 96/30037 (PCT/US96/02349), both of which are incorporated herein by reference).

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ischemia/reperfusion injury (as described in U.S. Patent No. 5,578,568, incorporated herein by reference), and liver resection (as described in co-pending U.S. Application Serial No. 08/582,230 filed March 16, 1998 which is a continued prosecution application of the same serial no. filed January 3, 1996, which is in turn a continuation of U.S. Application Serial No. 08/318,357 filed October 5, 1994, which is in turn a continuation-in-part of U.S. Application Serial No. 08/132,510 filed October 5, 1993, and corresponding International Publication No. WO 95/10297 (PCT/US94/11404), all of which are incorporated herein by reference).

BPI protein products are also known to neutralize the anti-coagulant activity of exogenous heparin, as described in U.S. Patent No. 5,348,942, incorporated herein by reference, as well as to be useful for treating chronic inflammatory diseases such as rheumatoid and reactive arthritis, as described in U.S. Patent No. 5,639,727, incorporated herein by reference, and for inhibiting angiogenesis and for treating angiogenesis-associated disorders including malignant tumors, ocular retinopathy and endometriosis, as described in co-pending U.S. Application Serial Nos. 08/435,855, 08/466,624 and 08/466,826, and corresponding International Publication No. WO 94/20128 (PCT/US94/02401), all of which are incorporated herein by reference.

BPI protein products are also known for use in antithrombotic methods, as described in U.S. Patent No. 5,741,779 and corresponding International Publication No. WO97/42967 (PCT/US97/08017), which are incorporated herein by reference.

U.S. Patent Nos. 5,420,019 and 5,674,834 and corresponding International Publication No. WO94/18323 (PCT/US94/01235), all of which are incorporated herein by reference, disclose that the replacement of the cysteine residue at amino acid position 132 or 135 with another amino acid renders the resulting BPI polypeptide resistant to dimerization and cysteine adduct formation. It also discloses that terminating the N-terminal BPI fragment at BPI amino acid position 193 resulted in an expression product with reduced carboxy-terminal heterogeneity.

Because of the multiplicity of valuable activities and uses of BPI protein products as exemplified above, a need continues to exist for new products with structures based on or mimicking a BPI protein product and having one or more of the activities and/or uses of BPI protein products, including use as anti-infective products, including antimicrobial agents (e.g., gram-negative bacteria [U.S. Patent

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Nos. 5,198,541 and 5,523,288; WO95/08344 (PCT/US94/11225)] and gram-positive bacteria [U.S. Patent No. 5,578,572; WO95/19180 (PCT/US95/00656)], fungi [U.S. Patent No. 5,627,153; WO95/19179 (PCT/US95/00498)], mycobacteria [EP0690721; WO94/20129 (PCT/US94/02463)] and chlamydia [WO96/01647 (PCT/US95/08624)] and endotoxin binding/neutralizing agents [WO95/019784 (PCT/US95/01151)], and as heparin binding/neutralizing products [U.S. Patent Nos. 5,348,942 and 5,639,727; WO94/20128 (PCT/US94/02401)], including for the neutralization of exogenously administered heparin, inhibition of angiogenesis (normal or pathological) for the treatment of chronic inflammatory disease states, and anticoagulant and thrombolytic agents for the treatment of thrombotic disorders [PCT/US97/08017]. All of the above-listed references regarding biological or functional activities of BPI, as well as therapeutic and diagnostic uses of BPI, are hereby incorporated by reference. One avenue of investigation toward solving the problem of new products based on BPI and fulfilling this need is the determination of the crystal structure of a BPI protein product.

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Summary of the Invention

The present invention solves the above problem and fulfills the need for designating and making new and useful products based on BPI. It is an object of this invention to solve the three-dimensional structure of BPI and thereby provide the atomic coordinates (i.e., structure coordinates) of BPI from the analysis of x-ray diffraction patterns of sufficiently high resolution to be useful for determining the three-dimensional protein structure.

It is an object of this invention to provide methods of expressing, purifying and crystallizing bactericidal/permeability-increasing protein (BPI) products, and thereby provide crystallized BPI proteins.

It is an object of this invention to provide the use of the structure coordinates of a BPI crystal to allow the design of compounds for mimicking a BPI protein product to reveal the atomic details of ligand binding sites of BPI (e.g. lipid-like or heparin-like molecules).

It is an object of this invention to provide use of the structure coordinates of a BPI crystal as described herein to solve the crystal structure of a crystal of a different BPI protein or fragment, analog or variant thereof, or a crystal of a related

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protein, including a BPI-related lipid transfer protein or a fragment, analog or variant thereof.

It is an object of this invention to provide mutants of BPI or fragments, analogs, or variants thereof characterized by one or more different properties as compared with wild-type BPI. These properties include altered surface charge, altered lipid binding pockets, altered specificity or higher activity. BPI mutants are useful to identify those amino acids that are most important for the lipid and heparin binding activity and other biological activities of BPI. This information, in turn, allows the design of new structures with one or more different properties based on BPI.

It is an object of this invention to provide the use of the structure coordinates and atomic details of BPI as described herein or its fragments, analogs or variants (including mutants or co-complexes) or of a BPI-related lipid transfer protein or its fragments, analogs or variants (including mutants or co-complexes) to design, evaluate computationally, synthesize and use new structures based on BPI with desirable properties of BPI, such as physical and pharmacological properties.

X-ray diffraction patterns of a related protein can be analyzed directly to provide the three-dimensional structure (if of sufficiently high resolution), however, the atomic coordinates for the crystallized BPI, as provided herein, can be used for structure determination. The x-ray diffraction patterns obtained by methods of the present invention, and provided on computer readable media, are used to provide electron density maps. The amino acid sequence is also useful for three-dimensional structure determination. The data is then used in combination with phase determination (e.g. using multiple isomorphous replacement (MIR) molecular replacement techniques) to generate electron density maps of BPI, using a suitable computer system. The electron density maps, provided by analysis of either the x-ray diffraction patterns or working backwards from the atomic coordinates, provided herein, are then fitted using suitable computer algorithms to generate secondary, tertiary and/or quaternary structures and/or domains of BPI, which structures and/or domains are then used to provide an overall three-dimensional structure, as well as binding and/or active sites of BPI.

It is also an object of this invention to specifically provide for the use of three-dimensional modeling of BPI and other members of the BPI protein family using the coordinates from the x-ray diffraction patterns. The coordinates and amino acid sequences are entered into one or more computer programs for molecular modeling. Such molecular modeling programs generate atomic coordinates that reflect the secondary, tertiary and/or quaternary structures of the protein which contribute to its overall three-dimensional structure and provide information related to binding and/or active sites of the protein.

It is a further object of this invention to specifically provide for the use of similar molecular modeling for rational drug design (RDD) of mimetics and ligands of BPI and other members of the BPI protein family. The drug design paradigm uses computer modeling programs to determine potential mimetics and ligands which are expected to interact with sites on the protein. The potential mimetics or ligands are then screened for activity and/or binding. For BPI-related mimetics or ligands, screening methods can be selected from assays for at least one biological activity of BPI, e.g., anti-microbial, LPS- binding/neutralizing, heparin binding/neutralizing, and/or anti-thrombotic activities, according to known method steps. Similarly for LBP-, CETP- or PLTP-related mimetics or ligands, such screening methods can be selected from assays for at least one biological activity of LBP, CETP or PLTP, according to known method steps. The resulting mimetics or ligands are then provided by methods of the present invention and are useful for treating, inhibiting or preventing BPI-modulated diseases (or LBP-, CETP- and PLTP-modulated) in animals, including humans.

Thus, as described herein, the present invention provides use of atomic coordinates of a BPI protein, or fragment, analog or variant thereof, to model a BPI protein or a related protein, including a BPI-related lipid transfer protein, such as LBP, CETP or PLTP, or fragment, analog or variant thereof.

The present invention also provides use of atomic coordinates of a BPI protein wherein the BPI protein comprises a binding site characterized by amino acid residues of at least one binding pocket as defined in Table 3, and/or wherein the BPI protein comprises a binding site characterized by at least one amino acid sequence, or variant of the sequence, selected from positions about 17 to about 45, positions about 65 to

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about 99 or positions about 142 to about 169 of BPI, or alternatively, a binding site characterized by at least one amino acid sequence, or variant of the sequence, selected from positions about 36 to about 54, positions about 84 to about 109 or positions about 142 to about 164 of BPI.

The present invention provides use of atomic coordinates of a BPI protein to computationally design a chemical compound for mimicking a BPI protein, or fragment, analog or variant thereof, or a BPI-related lipid transfer protein, or fragment, analog or variant thereof, including, for example, lipopolysaccharide-binding protein (LBP), cholesteryl ester transferase protein (CETP) or phospholipid transfer protein (PLTP), or fragment, analog, or variant thereof.

The present invention also provides use of atomic coordinates of BPI protein to design a chemical compound capable of associating with a BPI-related lipid binding protein, or fragment, analog or variant thereof, including, for example, bactericidal/permeability-increasing protein (BPI), lipopolysaccharide-binding protein (LBP), cholesteryl ester transferase protein (CETP) or phospholipid transfer protein (PLTP), or fragment, analog or variant thereof.

The present invention provides use of atomic coordinates of a BPI protein to design a model of ligands in an active site of a lipid binding protein, including, for example, BPI protein, lipopolysaccharide-binding protein (LBP), cholesteryl ester transferase protein (CETP) or phospholipid transfer protein (PLTP), or fragment, analog or variant thereof.

The present invention provides use of atomic coordinates of a bactericidal/permeability-increasing ("BPI") protein, to design compounds with at least one activity selected from the group consisting of antibacterial, antifungal, antimycobacterial, antichlamydial, antiprotozoan, heparin-binding, endotoxin-binding, heparin-neutralizing, endotoxin-neutralizing, inhibition of tumor and endothelial cell proliferation, inhibition of angiogenesis, anti-inflammatory, anticoagulant and antithrombolytic. The coordinates disclosed herein are suitable for all of the aforementioned uses of atomic coordinates.

The present invention provides a method of three-dimensional modeling of a BPI protein or BPI-related lipid transfer protein comprising the steps of: (a) providing three-dimensional atomic coordinates derived from X-ray diffraction measurements

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of a BPI protein in a computer readable format; (b) inputting the data from step (a) into a computer with appropriate software programs; and (c) generating a three-dimensional structural representation of the BPI protein or BPI-related lipid transfer protein suitable for visualization and further computational manipulation; particularly wherein the BPI protein comprises a binding site characterized by amino acid residues of at least one binding pocket as defined in Table 3, or wherein the BPI protein comprises a binding site characterized by at least one amino acid sequence, or variant of the sequence, selected from positions about 17 to about 45, positions about 65 to about 99 or positions about 142 to about 169 of BPI or wherein the BPI protein comprises a binding site characterized by amino acid residues of at least one binding pocket as defined in Table 3 and a binding site characterized by at least one amino acid sequence, or variant of the sequence, selected from positions about 36 to about 54, positions about 84 to about 109 or positions about 142 to about 164 of BPI, or alternatively from about positions 36 to about 54, from about positions 84 to about 109, or about positions 142 to about 164.

The present invention provides a method for providing an atomic model of a BPI protein, BPI-related lipid binding protein, or fragment, analog or variant thereof, comprising (a) providing a computer readable medium having stored thereon atomic coordinate/x-ray diffraction data of a BPI protein, or fragment, analog or variant thereof, in crystalline form, the data sufficient to model the three-dimensional structure of the BPI protein, or fragment, analog or variant thereof; (b) analyzing, on a computer using at least one subroutine executed in said computer, atomic coordinate/x-ray diffraction data from (a) to provide atomic coordinate data output defining an atomic model of said BPI protein, BPI-related binding lipid protein or fragment, analog or variant thereof, said analyzing utilizing at least one computing algorithm selected from the group consisting of data processing and reduction, autoindexing, intensity scaling, intensity merging, amplitude conversion, truncation, molecular replacement, molecular alignment, molecular refinement, electron density map calculation, electron density modification, electron map visualization, model building, rigid body refinement, positional refinement; and (c) obtaining atomic coordinate data defining the three-dimensional structure of at least one of said BPI protein, BPI-related lipid binding protein, or fragment, analog or variant thereof;

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particularly wherein said computer readable medium further has stored thereon data corresponding to a nucleic acid sequence or an amino acid sequence data comprising at least one structural domain or functional domain of a BPI, LBP, CETP or PLTP corresponding to at least one BPI or mutant primary sequence or fragment, analog or variant thereof; and wherein said analyzing step further comprises analyzing said sequence data.

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The present invention provides a computer-based system for providing atomic model data of the three-dimensional structure of a BPI protein, BPI-related lipid binding protein or fragment, analog or variant thereof, comprising the following elements: (a) at least one computer readable medium (CRM) having stored thereon atomic coordinate/x-ray diffraction data of a BPI protein, or fragment, analog or variant thereof; (b) at least one computing subroutine that, when executed in a computer, causes the computer to analyze atomic coordinate/x-ray diffraction data from (a) to provide atomic coordinate data output defining an atomic model of a BPI protein, BPI-related lipid binding protein or fragment, analog or variant thereof, said analyzing utilizing at least one computing subroutine selected from the group consisting of data processing and reduction, auto-indexing, intensity scaling, intensity merging, amplitude conversion, truncation, molecular replacement, molecular alignment, molecular refinement, electron density map calculation, electron density modification, electron map visualization, model building, rigid body refinement, positional refinement; and (c) retrieval means for obtaining atomic coordinate output data substantially defining the three-dimensional structure of said BPI protein, BPIrelated lipid binding protein or fragment, analog or variant thereof.

The present invention provides a method for providing a computer atomic model of a ligand of a BPI protein, BPI-related lipid binding protein, or fragment, analog or variant thereof, comprising: (a) providing a computer readable medium (CRM) having stored thereon atomic coordinate data of a BPI protein, or fragment, analog, or variant thereof; (b) providing a CRM having stored thereon atomic coordinate data sufficient to generate atomic models of potential ligands of said BPI protein, BPI-related lipid binding protein, or fragment, analog, or variant thereof; (c) analyzing on a computer, using at least one subroutine executed in said computer, the atomic coordinate data from (a) and ligand data from (b), to determine binding sites

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of BPI protein, BPI-related lipid binding protein, or fragment, analog, or variant thereof, and to provide atomic coordinate data defining an atomic model of at least one ligand of said BPI protein, BPI-related lipid binding protein, or fragment, analog or variant thereof, said analyzing utilizing computing subroutines selected from the group consisting of data processing and reduction, auto-indexing, intensity scaling, intensity merging, amplitude conversion, truncation, molecular replacement, molecular alignment, molecular refinement, electron density map calculation, electron density modification, electron map visualization, model building, rigid body refinement, positional refinement; and (d) obtaining atomic coordinate model output data defining the three-dimensional structure of said at least one ligand of said BPI protein, BPI-related lipid binding protein, or fragment, analog, or variant thereof.

The present invention provides a computer-based system for providing an atomic model of at least one ligand of a BPI protein, BPI-related lipid binding protein, or fragment, analog or variant thereof, comprising the following elements: (a) a computer readable medium (CRM) having stored thereon atomic coordinate data of a BPI protein, fragment, analog or variant thereof; (b) a CRM having stored thereon atomic coordinate data sufficient to generate atomic models of potential ligands of a BPI protein, BPI-related lipid binding protein, or fragment, analog or variant thereof; (c) at least one computing subroutine for analyzing on a computer, the atomic coordinate data from (a) and (b), to determine binding sites of BPI protein, BPI-related lipid binding protein. or fragment, analog, or variant thereof, and to provide data output defining an atomic model of at least one potential ligand of BPI protein, BPI-related lipid binding protein, or fragment, analog, or variant thereof, said analyzing utilizing at least one computing subroutine selected from the group consisting of data processing and reduction, auto-indexing, intensity scaling, intensity merging, amplitude conversion, truncation, molecular replacement, molecular alignment, molecular refinement, electron density map calculation, electron density modification, electron map visualization, model building, rigid body refinement, positional refinement; and (d) retrieval means for obtaining atomic coordinate data of said at least one ligand of a BPI protein, BPI-related lipid binding protein, or fragment, analog or variant thereof.

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said at least one ligand of a BPI protein, BPI-related lipid binding protein, or fragment, analog or variant thereof.

Other objects of the invention will be apparent to one of ordinary skill in the art from the following detailed description and examples relating to the present invention.

Brief Description of the Figures

- Fig. 1 (A) A ribbon diagram of residues 1-456 of BPI illustrating its boomerang shape. The NH₂-terminal domain is shown; the COOH-terminal domain and the two phosphatidylcholine molecules are shown. The linker is also shown, and the disulfide bond is shown as a ball-and-stick model. (B) View after rotating (A) 70° about the long axis of the molecule. Figure produced with MOLSCRIPT [P. Krauliz, *J. Appl. Cryst.*, 24:926 (1991)] and RASTER3D [E. A. Merrit and M. E. P. Murphy, *Acta Crystallogr.*, D50:889 (1994); D. J. Bacon and W. F. Anderson, *J. Mo. Graphics*, 6:219 (1988)].
- Fig. 2 (A) Schematic drawing of the novel BPI domain fold, shown in same orientation as the NH₂-terminal domain in Fig. 1B (B) Superposition of the NH₂- and COOH-terminal domains of BPI showing the overall topological similarity. Residues 1 to 230 and 250 to 456 are shown. The NH₂-terminal domain is in the same orientation as Fig. 1A.
- Fig. 3 Electron density of the final 2.8 Å MIR map contoured at 1.0 σ and superimposed on the refined model. The area shown is in the lipid binding pocket of the NH₂-terminal domain of BPI. The phosphatidylcholine and the surrounding protein atoms are shown.
- Fig. 4 (A) The covalent structure of phosphatidylcholine and the lipid A region of LPS from E. coli and S. typhimurium. Phosphate groups are indicated by P. Adapted with changes from [C. R. H. Raetz, Annu. Rev. Biochem, 59:129 (1990)].

 (B) Slice through the interior of BPI showing the lipid binding pocket in the NH₂-terminal domain. The solvent accessible surface of the protein was calculated without lipid present, the interior of the protein and the phosphatidylcholine are shown.
- 30 Protein residues are shown as ball-and-stick.

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Figure produced with MSP [M. L. Connolly, Science, 221:709 (1983); M. L. Connolly, J. Am. Chem. Soc., 107:1118 (1985)].

Figs. 5 (A) and 5 (B) The amino acid sequences of human BPI, LBP, PLTP, and CETP. The alignment was performed with CLUSTAL [D. G. Higgins and P. M. Sharp, Gene, 73:237 (1989)] using all eleven known protein sequences from 5 mammals [R. R. Schuman, et al., Science, 249:1429 (1990); D. Drayna et al., Nature, 327:632 (1987); R. Day et al., J. Biol. Chem., 269:9388 (1994); S.R. Leong and T. Camerato, Nucleic Acids Res., 18:3052 (1990); M. Nagashima, J. W. McLean, R. M. Lawn, J. Lipid Res., 29:1643 (1988); M. E. Pape, E. F. Rehber, K. 10 R. Marotti, G. W. Melchior, Artheriosclerosis 11:1759 (1991); G. Su et al., J. Immunol., 153:743 (1994); P. W. Gray et al., J. Biol. Chem. 264: 9505 (1989); Albers et al., Biochem. Biophys. Acta, 1258:27 (1995); X. C. Jiang et al., Biochemistry, 34:7258 (1995); L. B. Agellon et al., Biochemistry, 29:1372 (1990); X. C. Jiang et al., J. Biol. Chem., 266:4631 (1991)] but only the four human 15 sequences are shown. Residues that are completely conserved in all proteins are indicated below the sequence *; those which are highly conserved are indicated by •. The secondary structure of BPI is indicated above the sequences. The β strands are indicated by arrows; strands which make up the central β sheet are shown with gray arrows. Because of the β bulges and pronounced twisting, some of the β strands have one or more residues that do not show classical H-bonding patterns or $\Phi\Psi$ 20 angles; these breaks are indicated by $\hat{}$ above the strands. The α helices are shown as cylinders, and one-residue breaks in helices B and B' are indicated with a vertical dashed line. The horizontal dashed line indicates the linker region. Peptides from BPI and LBP with the highest LPS-binding activity (Little, et al., J. Biol. Chem. 268: 1865 (1994); Taylor et al., J. Biol. Chem. 270: 17934 (1995)) are in bold italics. 25 The disulfide bond is indicated by S-S. Residues with atoms within 4 Å of the NH₂terminal lipid are highlighted with gray shading; residues within 4 Å of the COOHterminal lipid are shown with white letters in black boxes.

Fig. 6 Block diagram of a computer system 102 that can be used to implement the present invention. The computer system 102 includes a processor 106 connected to

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a bus 104. Also connected to the bus 104 are a main memory 108 (preferably implemented as random access memory, RAM) and a variety of secondary storage memory 110, such as a hard drive 112, a removable medium storage device 114, a command device 118, and a visualization device, 120. Also included is a removable storage medium 116.

Detailed Description

The present invention provides methods for crystallizing a BPI protein product where the crystals diffract x-rays with sufficiently high resolution to allow determination of the three-dimensional structure of the BPI protein product, including atomic coordinates. The three-dimensional structure (e.g., as provided on computer readable media as described herein) is useful for rational drug design of BPI-related (and LBP-, CETP-, PLTP-related) mimetics and/or ligands. Specifically provided is a method for crystallizing a recombinant non-glycosylated human BPI analog holoprotein comprising a 456 amino acid sequence wherein the amino acid serine at position 351 has been changed to alanine. The three-dimensional structure is useful for modeling and/or synthesizing BPI-related mimetics or ligands. Such BPI-related mimetics or ligands are useful for treating, inhibiting or preventing BPI-modulated diseases.

The present invention thus includes methods of expressing, purifying and crystallizing a BPI protein product from suitable sources, such as eukaryotic cells or tissues. The present invention also provides crystallized BPI protein products by these methods. The crystallized BPI is analyzed by x-ray diffraction techniques to obtain high resolution diffraction patterns and atomic coordinates that are suitable for molecular modeling.

As used herein, "BPI protein product" or "BPI protein" includes naturally and recombinantly produced BPI protein; natural, synthetic, and recombinant biologically active polypeptide fragments of BPI protein; biologically active polypeptide variants of BPI protein or fragments thereof, including hybrid fusion proteins and dimers; biologically active polypeptide analogs of BPI protein or fragments or variants thereof, including cysteine-substituted analogs; and BPI-derived peptides. The BPI protein products for therapeutic or diagnostic uses may be generated and/or isolated by any means known in the art. U.S. Patent No. 5,198,541, the disclosure of which

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is incorporated herein by reference, discloses recombinant genes encoding and methods for expression of BPI proteins including recombinant BPI holoprotein, referred to as rBPI (also referred to as rBPI₅₅ or simply rBPI₅₀) and recombinant fragments of BPI. U.S. Patent Application Ser. No. 07/885,501, now abandoned, and a continuation-in-part thereof, U.S. Patent Application Ser. No. 08/072,063, filed May 19, 1993, issued as U.S. Patent 5,439,807 on August 8, 1995 and corresponding PCT Application No. 93/04752 filed May 19, 1993, which are all incorporated herein by reference, disclose novel methods for the purification of recombinant BPI protein products expressed in and secreted from genetically transformed mammalian host cells in culture and discloses how one may produce large quantities of recombinant BPI products suitable for incorporation into stable, homogeneous pharmaceutical preparations.

Biologically active fragments of BPI (BPI fragments) include biologically active molecules that have the same or similar amino acid sequence as a natural human BPI holoprotein, except that the fragment molecule lacks amino-terminal amino acids, internal amino acids, and/or carboxy-terminal amino acids of the holoprotein. Nonlimiting examples of such fragments include a N-terminal fragment of natural human BPI of approximately 25 kD, described in Ooi et al., J. Exp. Med., 174:649 (1991), and the recombinant expression product of DNA encoding Nterminal amino acids from 1 to about 193 or 199 of natural human BPI, described in Gazzano-Santoro et al., Infect. Immun. 60:4754-4761 (1992), and referred to as rBPI₂₃. In that publication, an expression vector was used as a source of DNA encoding a recombinant expression product (rBPI₂₁) having the 31-residue signal sequence and the first 199 amino acids of the N-terminus of the mature human BPI, as set out in Figure 1 of Gray et al., supra, except that valine at position 151 is specified by GTG rather than GTC and residue 185 is glutamic acid (specified by GAG) rather than lysine (specified by AAG). Recombinant holoprotein (rBPI) has also been produced having the sequence (SEQ ID NOS: 1 and 2) set out in Figure 1 of Gray et al., supra, with the exceptions noted for rBPI₂₃ and with the exception that residue 417 is alanine (specified by GCT) rather than valine (specified by GTT). Other examples include dimeric forms of BPI fragments, as described in U.S. Patent No. 5,447,913, and corresponding PCT Application No. PCT/US95/03125, the

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disclosures of which are incorporated herein by reference. Preferred dimeric products include dimeric BPI protein products wherein the monomers are aminoterminal BPI fragments having the N-terminal residues from about 1 to 175 to about 1 to 199 of BPI holoprotein. A particularly preferred dimeric product is the dimeric form of the BPI fragment having N-terminal residues 1 through 193, designated rBPI₄₂ dimer.

Biologically active variants of BPI (BPI variants) include but are not limited to recombinant hybrid fusion proteins, comprising BPI holoprotein or biologically active fragment thereof and at least a portion of at least one other polypeptide, and dimeric forms of BPI variants. Examples of such hybrid fusion proteins and dimeric forms are described by Theofan et al. in U.S. Patent Application Serial No. 07/885,911, now abandoned, and a continuation-in-part application thereof, U.S. Patent Application Serial No. 08/064,693 filed May 19, 1993, issued as U.S. Patent 5,643,570 on July 1, 1997 and corresponding PCT Application No. US93/04754 filed May 19, 1993, which are all incorporated herein by reference and include hybrid fusion proteins comprising, at the amino-terminal end, a BPI protein or a biologically active fragment thereof and, at the carboxy-terminal end, at least one constant domain of an immunoglobulin heavy chain or allelic variant thereof. Similarly configured hybrid fusion proteins involving part or all Lipopolysaccharide Binding Protein (LBP) are also contemplated for use in the present invention.

Biologically active analogs of BPI (BPI analogs) include but are not limited to BPI protein products wherein one or more amino acid residues have been replaced by a different amino acid. For example, U.S. Patent No. 5,420,019 and corresponding PCT Application No. US94/01235, filed February 2, 1994, the disclosures of which are incorporated herein by reference, discloses polypeptide analogs of BPI and BPI fragments wherein a cysteine residue is replaced by a different amino acid. A preferred BPI protein product described by this application is the expression product of DNA encoding from amino acid 1 to approximately 193 or 199 of the N-terminal amino acids of BPI holoprotein, but wherein the cysteine at residue number 132 is substituted with alanine and is designated rBPI₂₁Δcys or rBPI₂₁. Other examples include dimeric forms of BPI analogs; e.g. U.S. Patent Application Serial No. 08/212,132 filed March 11, 1994, issued as U.S. Patent 5,447,913 on

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September 5, 1995 and corresponding PCT Application No. PCT/US95/03125, the disclosures of which are incorporated herein by reference.

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Other BPI protein products useful according to the methods of the invention are peptides derived from or based on BPI produced by recombinant or synthetic means (BPI-derived peptides), such as those described in U.S. Patent Application Serial No. 08/504,841 filed July 20, 1995 and in PCT Application No. PCT/US94/10427 filed September 15, 1994, which corresponds to U.S. Patent Application Serial No. 08/306,473 filed September 15, 1994, issued as U.S. Patent 5.652,332 on July 29, 1997, and PCT Application No. US94/02465 filed March 11, 1994, which corresponds to U.S. Patent Application Serial No. 08/209,762, filed March 11, 1994, issued as U.S. Patent 5,733,872 on March 31, 1998, which is a continuation-in-part of U.S. Patent Application Serial No. 08/183,222, filed January 14, 1994, now abandoned, which is a continuation-in-part of U.S. Patent Application Ser. No. 08/093,202 filed July 15, 1993, now abandoned, (for which the corresponding international application is PCT Application No. US94/02401 filed March 11, 1994), which is a continuation-in-part of U.S. Patent Application Ser. No. 08/030,644 filed March 12, 1993, issued as U.S. Patent 5,348,942 on September 20, 1994, the disclosures of all of which are incorporated herein by reference.

Presently preferred BPI protein products include recombinantly-produced N-terminal fragments of BPI, especially those having a molecular weight of approximately between 21 to 25 kD such as rBPI₂₃ or rBPI₂₁, or dimeric forms of these N-terminal fragments (e.g., rBPI₄₂ dimer). Additionally, preferred BPI protein products include rBPI₅₀ and BPI-derived peptides.

The administration of BPI protein products is preferably accomplished with a pharmaceutical composition comprising a BPI protein product and a pharmaceutically acceptable diluent, adjuvant, or carrier. The BPI protein product may be administered without or in conjunction with known surfactants, other chemotherapeutic agents or additional known anti-microbial agents. One pharmaceutical composition containing BPI protein products (e.g., rBPI₅₀, rRBPI₂₃) comprises the BPI protein product at a concentration of 1 mg/ml in citrate buffered

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saline (5 or 20 mM citrate, 150 mM NaCl, pH 5.0) comprising 0.1% by weight of poloxamer 188 (Pluronic F-68, BASF Wyandotte, Parsippany, NJ) and 0.002% by weight of polysorbate 80 (Tween 80, ICI Americas Inc., Wilmington, DE). Another pharmaceutical composition containing BPI protein products (e.g., rBPI₂₁) comprises the BPI protein product at a concentration of 2 mg/mL in 5 mM citrate, 150 mM NaCl, 0.2% poloxamer 188 and 0.002% polysorbate 80. Such combinations are described in PCT Application No. US94/01239 filed February 2, 1994, which corresponds to U.S. Patent Application Ser. No. 08/190,869 filed February 2, 1994, issued as U.S. Patent 5,488,034 on January 30, 1996, and U.S. Patent Application Ser. No. 08/012,360 filed February 2, 1993, now abandoned, the disclosures of all of which are incorporated herein by reference. Additional formulations are provided in U.S. Patent Application Ser. Nos. 08/372,104, filed January 13, 1995, now abandoned, 08/530,599, filed September 19, 1995, now abandoned, and 08/586,133, filed January 12, 1996 and corresponding WO96/21436 (PCT/US96/01095).

The x-ray diffraction patterns of the invention are now discovered to be of sufficiently high resolution to be useful for three-dimensional modeling of a BPI. Preferably the resolution is in the range of 1.5 to 3.5 Å, preferably 1.5-3.0 Å and more preferably ≤ 2.6 Å.

Three-dimensional modeling is performed using the diffraction coordinates from these x-ray diffraction patterns. The coordinates are entered into one or more computer programs for molecular modeling, as known in the art. Such molecular modeling can utilize known x-ray diffraction molecular modeling algorithms or molecular modeling software to generate atomic coordinates corresponding to the three-dimensional structure of at least one BPI or a fragment thereof.

The entry of the coordinates of the x-ray diffraction patterns and the amino acid sequence into such programs results in the calculation of most probable secondary, tertiary and quaternary structures of the protein, including overall atomic coordinates of a BPI or a fragment thereof. These structures are combined and refined by additional calculations using such programs to determine the probable or actual three-dimensional structure of the BPI, including potential or actual active or binding sites of the protein.

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Such molecular modeling (and related) programs useful for rational drug design of ligands or mimetics, are also provided by the present invention. The drug design uses computer modeling programs which calculate how different molecules interact with the various sites of the BPI. This procedure determines potential ligands or mimetics of a BPI or at least one fragment thereof. The actual BPI-ligand complexes or mimetics are crystallized and analyzed using x-ray diffraction. The diffraction pattern coordinates are similarly used to calculate the three-dimensional interaction of a ligand and the BPI or a mimetic, in order to confirm that the ligand binds to, or changes the conformation of, a particular site on the BPI, or where the mimetic has a similar three-dimensional structure to that of a BPI or a fragment thereof.

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The potential ligands or mimetics are then screened for activity relating to a BPI. Such screening methods are selected from assays for at least one biological activity of the native BPI.

The resulting ligands or mimetics, provided by methods of the present invention, are useful for treating, screening or preventing bacterial infections in animals, such as mammals (including humans) and birds. Mimetics or ligands of a particular BPI will similarly react with other BPIs from other species, subgenera or genera of the BPI source organism.

Also provided are biologically active BPI proteins. A BPI protein is also provided as a crystallized protein suitable for x-ray diffraction analysis. The x-ray diffraction patterns obtained by the x-ray analysis are of moderately high to high resolution, e.g., 1.5-3.5 Å. The coordinates from these diffraction patterns are suitable and useful for three-dimensional modeling of the crystallized protein.

During the three-dimensional modeling of the BPI, these coordinates are entered with the BPI amino acid sequence into computer modeling programs to generate secondary, tertiary and quaternary structures of the BPI, as atomic coordinates. These structures together provide the three-dimensional structure of the BPI. The calculated and confirmed three-dimensional structure is then used for rational drug design of ligands or mimetics of the BPI or a fragment thereof.

The determination of the three-dimensional structure of a BPI protein thus has a broad-based utility. Significant sequence identity and conservation of important

structural elements is expected to exist among the BPI proteins of a particular species, subgenus, genus, or family. Therefore, the three-dimensional structure from one or a few BPI proteins can be used to identify therapeutics with one or more of the biological activities of BPI (and/or those of related proteins such as LBP, CETP and PLTP).

Determination of Protein Structures

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Different techniques give different and complementary information about protein structure. The primary structure is obtained by biochemical methods, either by direct determination of the amino acid sequence from the protein, or from the nucleotide sequence of the corresponding gene or cDNA. The quaternary structure of large proteins or aggregates can also be determined by electron microscopy. To obtain the secondary and tertiary structure, which requires detailed information about the arrangement of atoms within a protein, x-ray crystallography is preferred.

The first prerequisite for solving the three-dimensional structure of a protein by x-ray crystallography is a well-ordered crystal that will diffract x-rays strongly. The crystallographic method directs a beam of x-rays onto a regular, repeating array of many identical molecules so that the x-rays are diffracted from it in a pattern from which the structure of an individual molecule can be retrieved. Well-ordered crystals of globular protein molecules are large, spherical, or ellipsoidal objects with irregular surfaces, and crystals thereof contain large holes or channels that are formed between the individual molecules. These channels, which usually occupy more than half the volume of the crystal, are filled with disordered solvent molecules. The protein molecules are in contact with each other at only a few small regions. This is one reason why structures of proteins determined by x-ray crystallography are generally the same as those for the proteins in solution.

The formation of crystals is dependent on a number of different parameters, including pH, temperature, protein, concentration, the nature of the solvent and precipitant, as well as the presence of added ions or ligands to the protein. Many routine crystallization experiments may be needed to screen all these parameters for the few combinations that might give crystal suitable for x-ray diffraction analysis. Crystallization robots can automate and speed up the work of reproducibly setting up large number of crystallization experiments.

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A pure and homogeneous protein sample is important for successful crystallization. Proteins obtained from cloned genes in efficient expression vectors can be purified quickly to homogeneity in large quantities in a few purification steps. A protein to be crystallized is preferably at least 93-99% pure according to standard criteria of homogeneity. Crystals form when molecules are precipitated very slowly from supersaturated solutions. The most frequently used procedure for making protein crystals is the hanging-drop method, in which a drop of protein solution is brought very gradually to supersaturation by loss of water from the droplet to the larger reservoir that contains salt or polyethylene glycol solution.

Different crystal forms can be more or less well-ordered and hence give diffraction patterns of different quality. As a general rule, the more closely the protein molecules pack, and consequently the less water the crystals contain, the better is the diffraction pattern because the molecules are better ordered in the crystal.

X-rays are electromagnetic radiation at short wavelengths, emitted when electrons jump from a higher to a lower energy state. In conventional sources in the laboratory, x-rays are produced by high-voltage tubes in which a metal plate, the anode, is bombarded with accelerating electrons and thereby caused to emit x-rays of a specific wavelength, so-called monochromatic x-rays. The high voltage rapidly heats up the metal plate, which therefore has to be cooled. Efficient cooling is achieved by so-called rotating anode x-ray generators, where the metal plate revolves during the experiment so that different parts are heated up.

More powerful x-ray beams can be produced in synchrotron storage rings where electrons (or positrons) travel close to the speed of light. These particles emit very strong radiation at all wavelengths from short gamma rays to visible light. When used as an x-ray source, only radiation within a window of suitable wavelengths is channeled from the storage ring. Polychromatic x-ray beams are produced by having a broad window that allows through x-ray radiation with wavelengths of 0.2-3.5Å.

In diffraction experiments a narrow and parallel beam of x-rays is taken out from the x-ray source and directed onto the crystal to produce diffracted beams. The incident primary beam causes damage to both protein and solvent molecules. The crystal is, therefore, usually cooled to prolong its lifetime $(e.g., -220 \text{ to } -50^{\circ}\text{C})$. The

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primary beam must strike the crystal from many different directions to produce all possible diffraction spots, and so the crystal is rotated in the beam during the experiment.

The diffracted spots are recorded either on a film, the classical method, or by an electronic detector. The exposed film has to be measured and digitized by a scanning device, whereas electronic detectors feed the signals they detect directly in a digitized form into a computer. Electronic area detectors (an electronic film) significantly reduce the time required to collect and measure diffraction data.

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When the primary beam from an x-ray source strikes the crystal, some of the x-rays interact with the electrons on each atom and cause them to oscillate. The oscillating electrons serve as a new source of x-rays, which are emitted in almost all directions, referred to as scattering. When atoms (and hence their electrons) are arranged in a regular three-dimensional array, as in a crystal, the x-rays emitted from the oscillating electrons interfere with one another. In most cases, these x-rays, colliding from different directions, cancel each other out; those from certain directions, however, will add together to produce diffracted beams of radiation that can be recorded as a pattern on a photographic plate or detector.

The diffraction pattern obtained in an x-ray experiment is related to the crystal that caused the diffraction. X-rays that are reflected from adjacent planes travel different distances, and diffraction only occurs when the difference in distance is equal to the wavelength of the x-ray beam. This distance is dependent on the reflection angle, which is equal to the angle between the primary beam and the planes.

The relationship between the reflection angle (Θ), the distance between the planes (d), and the wavelength (λ) is given by Bragg's law: 2d sin $\Theta = \lambda$. This relation can be used to determine the size of the unit cell in the crystal. Briefly, the position on the film of the diffraction data relates each spot to a specific set of planes through the crystal. By using Bragg's law, these positions can be used to determine the size of the unit call.

Each atom in a crystal scatters x-rays in all directions, and only those that positively interfere with one another, according to Bragg's law, give rise to diffracted beams that can be recorded as a distinct diffraction spot above background. Each

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diffraction spot is the result of interference of all x-rays with the same diffraction angle emerging from all atoms. For example, for the protein crystal of myoglobin, each of the about 20,000 diffracted beams that have been measured contain scattered x-rays from each of the around 1500 atoms in the molecule. To extract information about individual atoms from such a system requires considerable computation. The mathematical tool that is used to handle such problems is called the Fourier transform.

Each diffracted beam, which is recorded as a spot on the film, is defined by three properties: the amplitude, which we can measure from the intensity of the spot; the wavelength, which is set by the x-ray source; and the phase, which is lost in x-ray experiments. All three properties are needed for all of the diffracted beams, in order to determine the position of the atoms giving rise to the diffracted beams.

For larger molecules, protein crystallographers have determined the phases in many cases using a method called multiple isomorphous replacement (MIR) (including heavy metal scattering), which requires the introduction of new x-ray scatterers into the unit cell of the crystal. These additions are usually heavy atoms (so that they make a significant contribution to the diffraction pattern), such that there should not be too many of them (so that their positions can be located); and they should not change the structure of the molecule or of the crystal cell, *i.e.*, the crystals should be isomorphous. Isomorphous replacement is usually done by diffusing different heavymetal complexes into the channels of the preformed protein crystals. The protein molecules expose side chains (such as SH groups) into these solvent channels that are able to bind heavy metals. It is also possible to replace endogenous light metals in metalloproteins with heavier ones, *e.g.*, zinc by mercury, or calcium by samarium.

Since such heavy metals contain many more electrons than the light atoms (H,N,C,O and S) of the protein, they scatter x-rays more strongly. All diffracted beams would therefore increase in intensity after heavy-metal substitution if all interference were positive. In fact, however, some interference is negative; consequently, following heavy-metal substitution, some spots measurably increase in intensity, others decrease, and many show no detectable difference.

Phase differences between diffracted spots can be determined from intensity changes following heavy-metal substitution. First, the intensity differences are used

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to deduce the positions of the heavy atoms in the crystal unit cell. Fourier summations of these intensity differences give maps of the vectors between the heavy atoms, the so-called Patterson maps. From these vector maps the atomic arrangement of the heavy atoms is deduced. From the positions of the heavy metals in the unit cell, one can calculate the amplitudes and phases of their contribution to the diffracted beams of protein crystals containing heavy metals.

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This knowledge is then used to find the phase of the contribution from the protein in the absence of the heavy-metal atoms. As both the phase and amplitude of the heavy metals and the amplitude of the protein alone is known, as well as the amplitude of the protein plus heavy metals (i.e., protein heavy-metal complex), one phase and three amplitudes are known. From this, the interference of the x-rays scattered by the heavy metals and protein can be calculated to see if it is constructive or destructive. The extent of positive or negative interference, with knowledge of the phase of the heavy metal, given an estimate of the phase of the protein. Because two different phase angles are determined and are equally good solutions, a second heavymetal complex can be used which also gives two possible phase angles. Only one of these will have the same value as one of the two previous phase angles; it therefore represents the correct phase angle. In practice, more than two different heavy-metal complexes are usually made in order to give a reasonably good phase determination for all reflections. Each individual phase estimate contains experimental errors arising from errors in the measured amplitudes. Furthermore, for many reflections, the intensity differences are too small to measure after one particular isomorphous replacement, and others can be tried.

The amplitudes and the phases of the diffraction data from the protein crystals are used to calculate an electron-density map of the repeating unit of the crystal. This map then has to be interpreted as a polypeptide chain with a particular amino acid sequence. The interpretation of the electron-density map is made more complex by several limitations of the data. First of all, the map itself contains errors, mainly due to errors in the phase angles. In addition, the quality of the map depends on the resolution of the diffraction data, which in turn depends on how well-ordered the crystals are. This directly influences the image that can be produced. The resolution

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is measured in Å units; the smaller this number is, the higher the resolution and therefore the greater the amount of detail that can be seen.

Building the initial model is a trial-and-error process. First, one has to decide how the polypeptide chain weaves its way through the electron-density map. The resulting chain trace constitutes a hypothesis, by which one tries to match the density of the side chains to the known sequence of the polypeptide. When a reasonable chain trace has finally been obtained, an initial model is built to give the best fit of the atoms to the electron density. Computer graphics are used both for chain tracing and for model building to present the data and manipulated the models.

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The initial model will contain some errors. Provided the protein crystals diffract to high enough resolution (e.g., better than 3.5 Å), most or substantially all of the errors can be removed by crystallographic refinement of the model using computer algorithms. In this process, the model is changed to minimize the difference between the experimentally observed diffraction amplitudes and those calculated for a hypothetical crystal containing the model (instead of the real molecule). This difference is expressed as an R factor (residual disagreement) which is 0.0 for exact agreement and about 0.59 for total disagreement.

In general, the R factor is preferably between 0.15 and 0.35 (such as less than about 0.24-0.28) for a well-determined protein structure. The residual difference is a consequence of errors and imperfections in the data. These derive from various sources, including slight variations in the conformation of the protein molecules, as well as inaccurate corrections both for the presence of solvent and for differences in the orientation of the microcrystals from which the crystal is built. This means that the final model represents an average of molecules that are slightly different both in conformation and orientation.

In refined structures at high resolution, there are usually no major errors in the orientation of individual residues, and the estimated errors in atomic positions are usually around 0.1-0.2 Å, provided the amino acid sequence is known. Hydrogen bonds, both within the protein and to bound ligands, can be identified with a high degree of confidence.

Those of skill in the art understand that a set of structure coordinates determined by X-ray crystallography is not without standard error. For the purpose

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of this invention, any set of structure coordinates for BPI-protein that have a root mean square deviation of protein backbone atoms (N, $C\alpha$, C and O) of less that 0.75Å when superimposed -- using backbone atoms -- on the structure coordinates listed in Table 4 shall be considered identical.

Most x-ray structures are determined to a resolution between 1.7 Å and 3.5 Å. Electron-density maps with this resolution range are preferably interpreted by fitting the known amino acid sequences into regions of electron density in which individual atoms are not resolved.

An amino acid sequence is preferred for accurate x-ray structure determination. Thus, recombinant DNA techniques have had a double impact on x-ray structural work. When a protein is cloned and overexpressed for structural studies, the amino acid sequence, necessary for the x-ray work, is also quickly obtained via the nucleotide sequence. Recombinant DNA techniques give us not only abundant supplies of rare proteins, but also their amino acid sequence as a bonus.

Overview of BPI Purification and Crystallization Methods

In general, a BPI protein is purified as described in Example 1. The resulting BPI is in sufficient purity and concentration for crystallization. The BPI is then isolated and assayed for biological activity and for lack of aggregation (which interferes with crystallization). The purified BPI preferably runs as a single band under reducing or nonreducing polyacrylamide gel electrophoresis (PAGE) (nonreducing is used to evaluate the presence of cysteine bridges).

The purified BPI is preferably crystallized using the hanging drop method under varying conditions of at least one of the following: pH, buffer type, buffer concentration, salt type, polymer type, polymer concentration, other precipitating agents and concentration of purified and cleaved BPI. See, e.g., the methods provided in a commercial kit, such as CRYSTAL SCREEN (Hampton Research, Riverside, CA). Differently sized and shaped crystals are tested for suitability for x-ray diffraction. Generally, larger crystals provide better crystallography than smaller crystals, and thicker crystals provide better diffraction than thinner crystals.

30 Purified BPIs

The results of the purification are optionally analyzed by polyacrylamide gel electrophoresis (PAGE) under reducing or non-reducing conditions. A single band

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is preferably obtained. With disulfide-containing BPIs, it is preferred that the analysis of the cleaved BPI be under non-reducing conditions to indicate whether the cleaved protein formed disulfide linked dimers. The amino acid sequence can also be determined according to known methods, or otherwise obtained, as this sequence is important in determining the three-dimensional structure of the cleaved protein (in combination with crystallographic analysis), as described herein, using molecular modeling techniques.

Before crystallization, the purified protein is tested for one or more of the known biological activities of a BPI protein.

It is preferred that the biological activity exceed the activity of the native protein. The preferred result indicates that the BPI protein retains its native structure, which is important for determining the three-dimensional crystal structure of the biologically active molecule. To identify the protease cleavage site, the purified and cleaved protein can be sequenced using known techniques. See, e.g., Murti et al., Proc. Natl. Acad. Sci. USA 90:1523-1525 (1993); Takimoto et al. (1992), infra, entirely incorporated herein by reference.

Protein Crystallization Methods

The hanging drop method is preferably used to crystallize the purified protein. See, e.g., Taylor et al., J. Mol. Biol. 226:1287-1290 (1992); Takimoto et al. (1992), infra; CRYSTAL SCREEN, Hampton Research.

A mixture of the purified protein and precipitant can include the following:

- pH (e.g., 4-9);
- buffer type (e.g., phosphate, cacodylate, acetates, imidazole, Tris HCl, sodium HEPES);
- buffer concentration (e.g., 10-200 mM);
 - salt type (e.g., calcium chloride, sodium citrate, magnesium chloride, ammonium acetate, ammonium sulfate, potassium phosphate, magnesium acetate, zinc acetate; calcium acetate)
 - polymer type and concentration: (e.g., polyethylene glycol (PEG)
 1-50%, average molecular weight 200-10,000);

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- other precipitating agents (salts: K, Na tartrate, NH₄SO₄, NaAc,
 LiSO₄, NaFormate, NaCitrate, MgFormate, NaPO₄, KPO₄ NH₄PO₄;
 organics: 2-propanol; non-volatile: 2-methyl-2,4-pentanediol); and
- concentration of purified BPI (e.g., 1.0-100 mg/ml).
- 5 See, e.g., CRYSTAL SCREEN, Hampton Research.

A non-limiting example of such crystallization conditions is the following:

- purified protein (e.g., approximately 3-4 mg/ml);
- H₂O;
- precipitant 10-14% Polyethylene glycol (PEG) 8000 buffered with 100
 mM cacodylate buffer and 200 mM of Mg acetate;
- at an overall pH of about 3.5-8.5.

The above mixtures are used and screened by varying at least one of pH, buffer type; buffer concentration, precipitating salt type or concentration, PEG type, PEG concentration, and protein concentration. Crystals ranging in size from 0.2-0.7 mm are formed in 1 -7 days. These crystals diffract x-rays to at least 3.5 Å resolution, such as 1.5 -3.5 Å, or any range of value therein, such as 1.5, 1.6, 1.7, 1.8, 1.9, 2.0, 2.1, 2.2, 2.3, 2.4, 2.5, 2.6, 2.7, 2.8, 2.9, or 3.0, with 3.0 Å or less being preferred.

Protein Crystals

20 Crystals appear after 1-4 days and grow to maximal size within a week. From one ten crystals are observed in one drop and crystal forms can occur, such as, but not limited to, bipyramidal, rhomboid, and cubic. Initial x-ray analyses indicate that such crystals diffract at moderately high to high resolution. When fewer crystals are produced in a drop, they can be much larger size, e.g., 0.4-0.9 mm.

25 X-ray Crystallography Methods and Molecular Modeling

The crystals so produced for BPI are x-ray analyzed using a suitable x-ray source. Diffraction patterns are obtained. Crystals are preferably stable for at least 10 hrs in the x-ray beam. Frozen crystals $(e.g., -220 \text{ to } -50^{\circ}\text{C})$ could also be used for longer x-ray exposures (e.g., 24-72 hrs), the crystals being relatively more stable to the x-rays in the frozen state. To collect the maximum number of useful reflections, multiple frames are optionally collected as the crystal is rotated in the x-ray beam, e.g., for 24-72 hrs. Larger crystals (>0.2 mm) are preferred, to increase

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the resolution of the x-ray diffraction. Alternatively, crystals may be analyzed using a synchrotron high energy x-ray source. Using frozen crystals, x-ray diffraction data is collected on crystals that diffract to a relatively high resolution of 3.5 Å or less, sufficient to solve the three-dimensional structure of BPI in considerable detail, as presented herein.

Native and/or derivative x-ray diffraction data with medium resolution is collected on area detectors mounted on rotating anode x-ray sources. The alternative program DENZO is preferably used for data processing and reduction (Sawyer et al., eds., Proceedings of CCP4 Study Weekend, pp. 56-62, SERC Darsbary Lab., UK (1993)).

The resolution is optionally improved using larger crystals, e.g., 0.2mm, making data collection more efficient, particularly for the determination of suitable heavy metal derivatives, such as Hg, Pt, Pb, Ba, Cd, and/or La derivatives.

The heavy metal derivatives are used to determine the phase, e.g., by the isomorphous replacement method. Heavy atom isomorphous derivatives of BPI are used for x-ray crystallography, where the structure is solved using one or several derivatives, which, (when combined) improves the overall figure of merit. Derivatives are identified through Patterson maps and/or cross-phase difference Fourier maps, e.g., using the CCP4 package (SERC Collaborative Computing Project No. 4, Daresbury Laboratory, UK, 1979).

Phases were also obtained or improved by optimization of the anomalous dispersion component of the x-ray scattering which can break the phase ambiguity which a single heavy atom derivative gives. In certain cases phase information may be obtained without the need of a native set of data, through the use of multiple wavelength with anomalous dispersion phasing (MAD phasing). The wavelength of the x-rays used may be selected at a synchrotron source to optimize this anomalous scattering. In this case data from a derivatised crystal or crystals is collected at typically three wavelengths, two of which are very close to the absorption edge of the heavy atom scatterer. One way of obtaining a suitable heavy atom derivatised crystal is to derivatise a known ligand of the protein.

The program MLPHARE (Wolf et al., eds., Isomorphous Replacement and Anomalous Scattering: Proceedings of CCP4 Study Weekend, pp. 80-86, SERC

Daresbury Lab., UK (1991)) is optionally used for refinement of the heavy atom parameters and the phases derived from them by comparing at least one of completeness (%), resolution (Å), R^r (%), heavy atom concentration (mM), soaking time, heavy atom sites, phasing power (acentric, centric) (See Table 1 as an analogous example from The Crystal Structure of diphtheria toxin, Choe et al., Nature 357: 216-222 (1992). Addition of heavy atom derivatives produce an MIR map with recognizable features.

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The initial phases are calculated to 3.2 Å, and then improved and extended to a higher resolution of 2.8 Å (e.g., \leq 3.0 Å) using solvent flattening, histogram matching and/or Sayre's equation in the program DM (Cowtan and Main, *Acta Crystallogr. D* 49:148-157 (1993)). The skeletonization of DM procedure is optionally used to improve connectivity in the bulk of the protein envelope. Both the MIR and density modified maps are optionally used in subsequent stages, to provide sufficient resolution and/or modeling of surface structures.

Skeletonized representations of electron density maps are then computed. These maps are automatically or manually edited using suitable software, e.g., the graphics package FRODO (Jones et al. (1991), infra) to give a continuous $C\alpha$ trace. The BPI sequence is then aligned to the trace. Initially pieces of idealized polypeptide backbone were placed into regions of the electron density map with obvious secondary structures (e.g., α -helix, β -sheet). After a polyalanine model was constructed for the protein, amino acid sidechains were added where density was present in the maps. The amino acid sequence of BPI was then examined for regions with distinct sidechain patterns (e.g., three consecutive aromatic rings). When a pattern in the sequence was found to match an area of the map, the correct sidechains were built onto the existing model. Eventually fragments containing recognizable sequence motifs were connected into a single chain, completing the tracing of the amino acid sequence into the maps.

X-ray diffraction data (e.g., to ≤ 3.0 Å) was collected on an RAXIS 11C area detector (e.g., a Mar imaging plate) mounted on a RIGAKU rotating anode or alternatively a synchrotron x-ray source, and processed using a suitable oscillation data reduction program (DENZO, Sawyer et al. eds., Proceedings of CCP4 Study Weekend, pp. 56-62, SERC Darsbary Lab., UK (1993). Cycles of simulated

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annealing against these data were refined using the program X-PLOR for molecular dynamics for R-factor refinement (X-PLOR, Brünger et al., J. Mol. Biol. 203:803-816 (1987)). This refinement was followed by manual rebuilding with FRODO using experimental and $2F_0$ - F_c maps. The model can be optionally further refined using a least-squares refinement program, such as TNT (Tronrud et al., Acta Crystallogr. A 43:489-501 (1987)).

One or more of the above modeling steps is performed to provide a molecular 3-D model of BPI. It is preferred that the BPI model has no residues in disallowed regions of the Ramachandran plot, and gives a positive 3D-1D profile (Luthy et al., Nature 356:83-85 (1992)), suggesting that all the residues are in acceptable environments (Kraulis (1991), infra).

Multiple isomorphous replacement phase determination was used for solving the three-dimensional structure of BPI. This structure is then used for rational drug design of BPI ligands or mimetics of at least one BPI bactericidal activity, or other biological activity important in inactivating bacterial toxicity, replication and/or infection.

Computer Related Embodiments

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An amino acid sequence of a BPI protein (or related protein such as LBP, CETP or PLTP) and/or x-ray diffraction data, useful for computer molecular modeling of BPI protein (or related protein such as LBP, CETP or PLTP) or a portion thereof, can be "provided" in a variety of mediums to facilitate use thereof. As used herein, provided refers to a manufacture, which contains, for example, a BPI amino acid sequence and/or atomic coordinate/x-ray diffraction data of the present invention, e.g., an amino acid sequence provided in Figure 5, a representative fragment thereof, or an amino acid sequence having at least 80-100% overall identity to an amino acid fragment of an amino acid sequence of Figure 5 or a variant thereof. Such a method provides the amino acid sequence and/or x-ray diffraction data in a form which allows a skilled artisan to analyze and molecular model the three-dimensional structure of a BPI-related protein, including a subdomain thereof.

In one application of this embodiment, BPI (or related protein such as LBP, CETP or PLTP), or at least one subdomain thereof, amino acid sequence and/or x-ray diffraction data of the present invention is recorded on computer readable medium.

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As used herein, "computer readable medium" refers to any medium which can be read and accessed directly by a computer. Such media include, but are not limited to: magnetic storage media, such as floppy discs, hard disc storage medium, and magnetic tape; optical storage media such as optical discs or CD-ROM; electrical storage media such as RAM and ROM; and hybrids of these categories such as magnetic/optical storage media. A skilled artisan can readily appreciate how any of the presently known computer readable mediums can be used to create a manufacture comprising computer readable medium having recorded thereon an amino acid sequence and/or x-ray diffraction data of the present invention.

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As used herein, "recorded" refers to a process for storing information on computer readable medium. A skilled artisan can readily adopt any of the presently know methods for recording information on computer readable medium to generate manufactures comprising an amino acid sequence and/or atomic coordinate/x-ray diffraction data information of the present invention.

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A variety of data storage structures are available to a skilled artisan for creating a computer readable medium having recorded thereon an amino acid sequence and/or atomic coordinate/x-ray diffraction data of the present invention. The choice of the data storage structure will generally be based on the means chosen to access the stored information. In addition, a variety of data processor programs and formats can be used to store the sequence and x-ray data information of the present invention on computer readable medium. The sequence information can be represented in a word processing text file, formatted in commercially-available software such as WordPerfect and MICROSOFT Word, or represented in the form of an ASCII file, stored in a database application, such as DB2, Sybase, Oracle, or the like. A skilled artisan can readily adapt any number of dataprocessor structuring formats (e.g. text file or database) in order to obtain computer readable medium having recorded thereon the information of the present invention.

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By providing computer readable medium having stored thereon a BPI or related sequence protein and/or atomic coordinates based on x-ray diffraction data, a skilled artisan can routinely access the sequence and atomic coordinate or x-ray diffraction data to model a BPI or related protein, a subdomain thereof, mimetic, or a ligand thereof. Computer algorithms are publicly and commercially available which

allow a skilled artisan to access this data provided in a computer readable medium and analyze it for molecular modeling and/or RDD. See, e.g., Biotechnology Software Directory, MaryAnn Liebert Publ., New York (1995).

The present invention further provides systems, particularly computer-based systems, which contain the sequence and/or diffraction data described herein. Such systems are designed to do structure determination and RDD for a BPI or related protein or at least one subdomain thereof. Non-limiting examples are microcomputer workstations available from Silicon Graphics Incorporated and Sun Microsystems running UNIX based, Windows NT or IBM OS/2 operating systems.

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As used herein, "a computer-based system" refers to the hardware means, software means, and data storage means used to analyze the sequence and/or x-ray diffraction data of the present invention. The minimum hardware means of the computer-based systems of the present invention comprises a central processing unit (CPU), input means, output means, and data storage means. A skilled artisan can readily appreciate which of the currently available computer-based systems are suitable for use in the present invention. A visualization device, such as a monitor, is optionally provided to visualize structure data.

As stated above, the computer-based systems of the present invention comprise a data storage means having stored therein a BPI or related protein or fragment sequence and/or atomic coordinate/x-ray diffraction data of the present invention and the necessary hardware means and software means for supporting and implementing an analysis means. As used herein, "data storage means" refers to memory which can store sequence or atomic coordinate/x-ray diffraction data of the present invention, or a memory access means which can access manufactures having recorded thereon the sequence or x-ray data of the present invention.

As used herein, "search means" or "analysis means" refers to one or more programs which are implemented on the computer-based system to compare a target sequence or target structural motif with the sequence or x-ray data stored within the data storage means. Search means are used to identify fragments or regions of a BPI or related protein which match a particular target sequence or target motif. A variety of known algorithms are disclosed publicly and a variety of commercially available software for conducting search means are and can be used in the computer-based

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systems of the present invention. A skilled artisan can readily recognize that any one of the available algorithms or implementing software packages for conducting computer analyses can be adapted for use in the present computer-based systems.

As used herein, "a target structural motif," or "target motif," refers to any rationally selected sequence or combination of sequences in which the sequence(s) are chosen based on a three-dimensional configuration or electron density map which is formed upon the folding of the target motif. There are a variety of target motifs known in the art. Protein target motifs include, but are not limited to, enzymic active sites, structural subdomains, epitopes, functional domains and signal sequences. A variety of structural formats for the input and output means can be used to input and output the information in the computer-based systems of the present invention.

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A variety of comparing means can be used to compare a target sequence or target motif with the data storage means to identify structural motifs or electron density maps derived in part from the atomic coordinate/x-ray diffraction data. A skilled artisan can readily recognize that any one of the publicly available computer modeling programs can be used as the search means for the computer-based systems of the present invention.

One application of this embodiment is provided in Figure 6. Figure 6 provides a block diagram of a computer system 102 that can be used to implement the present invention. The computer system 102 includes a processor 106 connected to a bus 104. Also connected to the bus 104 are a main memory 108 (preferably implemented as random access memory, RAM) and a variety of secondary storage memory 110, such as a hard drive 112, a removable medium storage device 114, a command device 118, and a visualization device, 120. Also included is a removable storage medium 116. The removable medium storage device 114 may represent, for example, a floppy disk drive, a CD-ROM drive, a magnetic tape drive, etc. A removable storage medium 116 (such as a floppy disk, a compact disk, a magnetic tape, etc.) containing control logic and/or data recorded therein may be inserted into the removable storage device 114. The computer system 102 includes appropriate software for reading the control logic and/or the data from the removable storage medium 116 once inserted in the removable medium storage device 114.

Amino acid, encoding nucleotide or other sequence and/or atomic coordinate/x-ray diffraction data of the present invention may be stored in a well known manner in the main memory 108, or any of the secondary storage devices 110, and/or a removable storage medium 116. Software for accessing and processing the amino acid sequence and/or atomic coordinate/x-ray diffraction data (such as search tools, comparing tools, etc.) reside in main memory 108 during execution. User commands are implemented through a command device 118, such as a keyboard. The visualization device 120 is optionally used to visualize the structure data.

Structure Determination

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One or more computational steps, computer programs and/or computer algorithms are used to provide a molecular 3-D model of BPI or related protein, using amino acid sequence data from Figure 5 (or fragments or variants thereof) and/or atomic coordinate/x-ray diffraction data. In x-ray crystallography, x-ray diffraction data and phases are combined to produce electron density maps in which the three-dimensional structure of a BPI protein is then built or modeled. MIR Phase determination was used for solving the three-dimensional structure of BPI. This structure can then be used for RDD of mimetics or ligands of a BPI or related protein and its associated biological activity, which is relevant to a protein modulated disease.

Electron density maps were calculated by X-PLOR or alternatively using such programs as those from the CCP4 computing package (SERC (UK) Collaborative Computing Project 4, Daresbury Laboratory, UK, 1979). If non-crystallographic symmetry axes are present, cycles of symmetry averaging can further be used, such as with the program RAVE (Kleywegt & Jones, Bailey et al., eds., First Map to Final Model, SERC Daresbury Laboratory, UK, pp 59-66 (1994)) and gradual model expansion. For map visualization and model building the program FRODO was used or alternatively, a program such as "O" (Jones (1991), infra) can be used.

Refinement and Model Validation

Density Modification and Map Interpretation

Rigid body and positional refinement can be carried out using a program such as X-PLOR (Brünger (1992), *infra*), e.g., with the stereochemical parameters of Engh and Huber (Acta Cryst. A47:392-400 (1991)). If the model at this stage in the

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averaged maps is still missing residues (e.g., at least 5-10 per subunit), some or all of the missing residues can be incorporated in the model during additional cycles of positional refinement and model building. The refinement procedure can start using data from lower resolution (e.g., 25-10Å to 10-3.0 Å and then be gradually extended to include higher resolution data from 12-6Å to 3.0-1.5 Å). B-values (also termed temperature factors) for individual atoms were refined once data between 2.9 and 1.5 A has been added. Subsequently waters were gradually added by manual inspection of electron density maps. Alternatively, a program such as ARP (Lamzin and Wilson, Acta Cryst. D49:129-147 (1993)) can be used to add crystallographic waters and as a tool to check for bad areas in the model. The programs PROCHECK (Lackowski et al., J. Appl. Cryst. 26:283-291 (1993)), WHATIF (Vriend, J. Mol. Graph. 8:52-56 (1990)), PROFILE 3D (Lüthy et al., Nature 356:83-85 (1992)), and ERRAT (Colovos & Yeates Protein Science, 2:1511-19 (1993)) as well as the geometrical analysis generated by X-PLOR were used to check the structure for errors. Anisotropic scaling between F_{obs} and F_{calc} was applied after careful assessment of the quality and completeness of the data.

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The program DSSP was used to assign the secondary structure elements (Kabsch and Sander, *Biopolymers*, 22:2577-2637 (1983)). A program such as SUPPOS (from the BIOMOL crystallographic computing package) can be used for some or all of the least-squares superpositions of various models and parts of models. The program ALIGN (Cohen *J. Mol. Biol.*, 190: 593-604 (1986)) was used to superimpose N- and C-terminal domains of BPI. Solvent accessible surfaces and electrostatic potentials can be calculated using such programs as GRASP (Nicholls *et al.* (1991), *infra*).

The structure of BPI from different organisms and the related proteins LBP, CETP and PLTP can thus be solved with the molecular replacement procedure such as by using X-PLOR (Brünger (1992), *infra*). A partial search model for a portion or all of these proteins can be constructed using the structures of BPI. The rotation and translation function can be used to yield orientations and positions for these models. Symmetry averaging can also be done using the RAVE program and model expansion can also be used to add missing residues resulting in a model with 95-99.9% of the total number of residues. The model can be refined in a program such

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as X-PLOR (Brünger (1992), supra), to a suitable crystallographic R_{factor} . The model data is then saved on computer readable medium for use in further analysis, such as rational drug design.

Rational Design of Mimetics or Ligands

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The determination of the crystal structure of a BPI protein, as described herein, provides a basis for the design of new and specific agents, including proteins or organic compounds.

Several approaches can be taken for the use of the crystal structure of a BPI in the rational design of protein or organic analogs having a relevant activity similar to that of a BPI or related protein. A computer-assisted, manual examination of a BPI potential binding site structure is optionally done. The use of software such as GRID — Goodford, J. Med. Chem. 28:849-857 (1985) a program that determines probable interaction sites between probes with various functional group characteristics and the protein surface — is used to analyze the surface sites to determine structures of similar inhibiting proteins or compounds. The GRID calculations, with suitable inhibiting groups on molecules (e.g., protonated primary amines) as the probe, are used to identify potential hotspots around accessible positions at suitable energy contour levels.

A diagnostic or therapeutic BPI or related protein modulating ligand of the present invention can be, but is not limited to, at least one selected from a lipid, a nucleic acid, a compound, a protein, an element, an antibody, a saccharide, an isotope, a carbohydrate, an imaging agent, a lipoprotein, a glycoprotein, an enzyme, a detectable probe, and antibody or fragment thereof, or any combination thereof, which can be detectably labeled as for labeling antibodies. Such labels include, but are not limited to, enzymatic labels, radioisotope or radioactive compounds or elements, fluorescent compounds or metals, chemiluminescent compounds and bioluminescent compounds. Alternatively, any other known diagnostic or therapeutic agent can be used in a method of the invention. Suitable compounds are then tested for activities of a BPI protein or BPI mimetic.

The program DOCK (Kuntz et al. J. Mol. Biol., 161:269-288 (1982)) may be used to analyze an active site or ligand binding site and suggest ligands with complementary steric properties. Several methodologies for searching three-

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dimensional databases to test pharmacophore hypotheses and select compounds for screening are available. These include the program CAVEAT (Bacon et al. *J. Mol Biol.*, 225: 849-858 (1992)) which uses databases of cyclic compounds which can act as "spacers" to connect any number of chemical fragments already positioned in the active site. This allows one skilled in the art to quickly generate hundreds of possible ways to connect the fragments already known or suspected to be necessary for tight binding. The program LUDI (Bohm et al. *J. Comput.-Aid. Mol. Des.*, 6:61-78 (1992)) can determine a list of interactions sites into which to place both hydrogen bonding and hydrophobic fragments. LUDI then uses a library of approx. 600 linkers to connect up to four different interaction sites into fragments. Then smaller "bridging" groups such as -CH2- and -COO- are used to connect these fragments. For example, for the enzyme DHFR, the placements of key functional groups in the well-known inhibitor methotrexate were reproduced by LUDI. See also, Rotstein and Murcko, *J. Med. Chem.*, 36:1700-1710 (1992)).

After preliminary experiments are done to determine the K_i of a ligand (e.g., a lipid ligand) by BPI (or related) protein to a BPI (or related) protein, mimetic or fragment, the time-dependent nature of the inhibition by the BPI or related protein (e.g., by the method of Henderson (Biochem. J. 127:321-333 (1972)) is determined.

For example, a lipid ligand and a BPI mimetic are pre-incubated in buffer. Reactions are initiated by the addition of detecting substrate. Aliquots are removed over a suitable time course and each quenched by addition into the aliquots of suitable quenching solution. The concentration of product are determined by known methods of detection. Plots of activity against time can be close to linear over the assay period, and are used to obtain values for the initial velocity in the presence (V_i) or absence (V_o) of, for example, a BPI mimetic. Error is present in both axes in a Henderson plot, making it inappropriate for standard regression analysis (Leatherbarrow, *Trends Biochem. Sci. 15*:455-458 (1990)). Therefore, K_i values are obtained from the data by fitting to a modified version of the Henderson equation for competitive inhibition:

$$Qr^2 + (E_t - Q - I_t)r - E_t = 0$$

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where (using the notation of Henderson (Biochem. J. 127:321-333 (1972)):

$$Q = K_t \left(\frac{A_t + K_a}{K_a} \right)$$

and

$$r = \frac{V_o}{V_i}$$

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This equation is solved for the positive root with the constraint that $Q = K_i((A_i + K_a) / K_a)$ using PROCNLIN from SAS (SAS Institute Inc., Cary, North Carolina, USA) which performs nonlinear regression using least-square techniques. The iterative method used is optionally the multivariate secant method, similar to the Gauss-Newton method, except that the derivatives in the Taylor series are estimated from the histogram of iterations rather than supplied analytically. A suitable convergence criterion is optionally used, e.g., where there is a change in loss function of less than 10^{-8} .

Once modulating compounds are found, crystallographic studies of co-complexes, for example, BPI mimetics complexed to a ligand are performed. As used herein, a co-complex refers to a BPI protein, fragment, analog or variant thereof in covalent or non-covalent association with a chemical entity or compound. As a non-limiting example, BPI crystals are soaked for 2 days in 0.01-100 mM inhibitor compound and x-ray diffraction data are collected on an area detector and/or an image plate detector (e.g., a Mar image plate detector) using a rotating anode x-ray source. Data are collected to as high a resolution as possible, e.g., $\leq 3.0 \text{ Å}$, and merged with a suitable R-factor on intensities. An atomic model of the mimetic is built into the difference Fourier map ($F_{\text{inhibitor complex}}$ - F_{native}). The model can be refined to convergence in a cycle of simulated annealing (Brünger (1987), infra) involving 10-100 cycles of energy refinement, 100-10,000 1-fs steps of room temperature dynamics and/or 10-100 more cycles of energy refinement. Harmonic restraints may be used

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for the atom refinement, except for atoms within a 10-15 Å radius of the inhibitor. An R-factor is calculated for the model as well as an r.m.s. deviation from the ideal bond lengths and angles.

Direct measurements of activity provide further confirmation that the modeled mimetic compounds are high-affinity inhibitors for the lipid ligands. Other suitable assays for biological activity known for BPI or related proteins may be used.

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Preferably, little or no change in the structure of the BPI or mimetic occurs in the electron density map described above. K_i values are determined by a previously described method (Henderson (1972), *infra*) to evaluate mimetic proteins or organic compounds.

Atomic coordinates of BPI proteins are useful in the generation of molecular models of related proteins and of BPI mimetics. The atomic coordinates generated from the solved three-dimensional structure of BPI disclosed herein may be utilized in combination with additional structural and/or physicochemical information, such as amino acid sequence data, x-ray diffraction data, combinations of x-ray diffraction data from multiple isomorphous replacement molecular replacement techniques, or other phase determination techniques. These combinations may be used to generate other three-dimensional coordinate data useful to generate secondary, tertiary and/or quaternary structures and/or domains of BPI or related proteins, including BPI-related lipid binding proteins, or their fragments, analogs, or variants. These alternate coordinate sets are useful to provide overall three-dimensional structure, as well as binding and or active sites of a BPI or related protein, including BPI-related lipid binding proteins, or their fragments, analogs, or variants. These alternate coordinate data sets are also useful in molecular modeling computer-based systems and methods for rational drug design of mimetics and ligands of BPI and other related proteins, including other BPI-related lipid binding proteins. Utilizing CLUSTAL (a multiple sequence alignment program in PC-Gene) and the Homology module (a structurebased homology modeling program in InsightII on a Silicon Graphics Incorporated workstation, molecular models (and the corresponding three-dimensional coordinates files) of lipopolysaccharide binding protein (LBP), cholesteryl ester transfer protein (CETP) and phospholipid transfer protein (PLTP) are generated. With these files, existing mutants are mapped and new ones designed.

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The results described herein demonstrate that tight-binding mimetics of a BPI or related protein, based on the crystal structure of BPI, are provided by the present invention. Demonstration of clinically relevant levels of a biological activity of the mimetic is also useful.

In evaluating mirnetics for biological activity in animal models (e.g., mouse, rat, rabbit, baboon) various oral and parenteral routes of administration are evaluated. Using this approach, it is expected that a biological activity occurs in suitable animal models, e.g., using the mimetics discovered by structure determination and x-ray crystallography.

Having now generally described the invention, the same will be more readily understood through reference to the following examples which are provided by way of illustration, and are not intended to be limiting of the present invention.

Example 1: Preparation and Purification for Crystallization Construction of Plasmids Containing BPI (S351A)

BPI contains a single N-linked glycosylation site at the asparagine at position 349 which was eliminated by genetic engineering of the DNA sequence of BPI as follows. For glycosylation to occur at this position, the asparagine must occur within the sequence Asn-X-Ser/Thr where X can be any amino acid, except proline. N-linked glycosylation can be eliminated by either changing the Asn to another amino acid such as glutamine or by changing the serine or threonine to an alternate amino acid. The latter strategy was used to construct vectors containing BPI with an alanine at position 351 instead of serine.

Construction of Plasmids for BPI Expression

The plasmid pIC108 containing a cDNA encoding BPI cloned in a T3T7 plasmid (Clontech, Palo Alto, CA) served as the starting point for the construction of a vector for expression of nonglycosylated rBPI in mammalian cells.

To allow insertion of BPI into an optimized mammalian expression vector, a unique XhoI site was first added to the 3' end of the BPI gene in pIC108. Two oligonucleotides were synthesized for this purpose: BPI-53 (5' ACT GGT TCC ATG GAG GTC AGC GCC 3') encoding amino acids 361 - 370 of BPI and BPI-54 (5' GAC AGA TCT CTC GAG TCA TTT ATA GAC AA 3') encoding the last four

amino acids of coding sequence, the stop codon (TGA), and incorporating an XhoI site immediately downstream of the stop codon. These oligonucleotides were used to PCR amplify a 280 bp fragment of the C-terminus of BPI and incorporate the XhoI site at the 3´ end of the gene. The amplified fragment was digested with NcoI and BgIII and ligated to a \sim 4100 bp NcoI-BamHI fragment from pIC108 to generate the plasmid pSS101.

Construction of Plasmid with BPI (S351A)

The glycosylation site was next removed by replacing the region from a unique *Xcm*I site to a unique *Sph*I site within the BPI gene in pSS101 with an annealed oligonucleotide that contained the codon (TCC) for the serine at amino acid position 351 changed to the codon (GCC) for alanine as shown below.

Wild type

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XcmI

...CCC AAC TCC TCC CTG GCT TCC CTC TTC CTG ATT GGC ATG CAC

15 ...GGG TTC AGG AGG GAC CGA AGG GAG AAG GAC TAA CCG TAC GTG

Pro Asn Ser Ser Leu Ala Ser Leu Phe Leu Ile Gly Met His

351

Nonglycosylated

XcmI

SphI

...ccc AAC TCC GCC CTG GCT TCC CTC TTC CTG ATT GGC ATG CAC
...ggg TTC AGG CGG GAC CGA AGG GAG AAG GAC TAA CCG TAC GTG

Pro Asn Ser Ala Leu Ala Ser Leu Phe Leu Ile Gly Met His
351

This step generated the plasmid pSS102.

To construct the vector, pING4322, for the expression of full length nonglycosylated holo BPI in mammalian cells, pSS102 was digested with *Bst*BI and *Xho*I and a 596 bp fragment, which included the modified BPI sequence, was purified and ligated to the large *Bst*BI-*Xho*I fragment from pING4147 which contains the gpt gene encoding resistance to mycophenolic acid, the human Ig enhancer, the human cytomegalovirus promoter (CMV) and the mouse light chain 3' untranslated region and is identical to the vector, pING4144 as described in U.S. Patent No. 5,420,019 and WO94/18323 (PCT/US94/01235) hereby incorporated by reference, except that it contains the codon for the native cysteine instead of an alanine at amino acid position 132 of the BPI gene.

Stable Transfection of Mammalian Cells for Expression of Nonglycosylated BPI

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Mammalian cells are preferred hosts for production of rBPI protein analogs as described herein. Such cells permit proper secretion, folding, and post-translational modification of expressed proteins. Presently preferred mammalian host cells for production of BPI proteins include cells of fibroblast origin, such as CHO-K1 cells (ATCC CCL61), CHO-DG44 cells (a dihydrofolate reductase [DHFR] minus mutant of CHO Toronto obtained from Dr. Lawrence Chasin, Columbia University), CHO-DXB-11 (a DHFR mutant of CHO-K1 obtained from Dr. Lawrence Chasin), Vero Cells (ATCC CRL81) and Baby Hamster Kidney (BHK) cells (ATCC CRL6281) and cells of lymphoid origin, such as the hybridoma Sp2/O-Ag14 (ATCC CRL1581) or the myeloma, NSO (ECACC No. 85110503).

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Transfection of mammalian cells can be accomplished by a variety of methods. Two of the most common approaches involve calcium phosphate precipitation of the expression vector DNA which is subsequently taken by the cells and electroporation, which causes the cells to take up the DNA through membrane pores created by the generation of a strong electric field. Selection for transfected cells is facilitated by the incorporation in the expression vector of a gene whose product allows the transfected cells to survive and grow under selective conditions. A number of such genes have been identified including, among others, the bacterial Tn5 neo gene, which encodes resistance to the antibiotic G418 and the Escherichia coli guanine phosphoribosyl transferase (gpt) gene, which encodes resistance to mycophenolic acid (MPA) in the presence of xanthine (Mulligan and Berg, Proc. Natl. Acad. Sci. 78:2072-2076 (1981)), the dihydrofolate reductase (DHFR) gene, which allows for growth of DHFR cells in the absence of nucleosides and gene amplification in the presence of increasing concentrations of methotrexate, the glutamine synthetase gene, which allows for growth of glutamine auxotrophs without glutamine and gene amplification in the presence of methionine sulfoximine and the Salmonella typhimurium hisD gene and the E. coli trpB gene (Hartman and Mulligan, Proc. Natl. Acad. Sci. 85: 8047-8051 (1988)), which allow growth in the presence of histidinol or without tryptophan (in the presence of indole), respectively. The availability of these selective markers provide significant flexibility for the generation of mammalian cell lines that express recombinant products, since they can be used either alone or in various combinations to provide cell lines with the highest possible productivity.

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Transfection of CHO-K1 Cells with pING4322

The CHO-K1 cell line was maintained in Ham's F12 medium plus 10% fetal bovine serum (FBS). Media were supplemented with glutamine/penicillin/streptomycin (Irvine Scientific, Irvine, California).

CHO-K1 cells were transfected by electroporation with 40 μ g of pING4322 DNA that was first digested with PvuI, phenol-chloroform extracted and ethanol precipitated. Following the electroporation, the cells were allowed to recover for 24 hours in non-selective Ham's F12 medium. The cells were then trypsinized, resuspended at a concentration of ~5 X 10⁴ cells/ml in Ham's F12 medium supplemented with MPA (25 μ g/mL) and xanthine (250 μ g/mL) and plated at ~ 10⁴ cells/well in 96 well plates. Untransfected CHO-K1 cells are unable to grow in this medium due to the inhibition of pyrimidine synthesis by the MPA. At ~ 2 weeks, colonies consisting of transfected cells were observed in the 96 well plates. Supernatants from wells containing single colonies were analyzed for the presence of BPI-reactive protein by anti-BPI ELISA using BPI₂₃ as a standard. In this assay, Immulon-II 96 well plates (Dynatech) were pre-coated with affinity purified rabbit anti-BPI₂₃ antiserum, followed by supernatant samples and detection was with affinity purified, biotinylated rabbit anti-BPI23 antiserum followed by peroxidase-labeled avidin. A total of 100 colonies were screened in this manner. The top isolates were transferred to 24 well plates and productivity was assessed as follows. Cells were grown to confluence in a 24 well plate in Ham's F12 medium supplemented with 10% FBS. Once the cells reached confluence, the Ham's F12 medium was removed and 2 ml of HB-CHO serum free medium (Irvine Scientific) plus 40 μL of S-Sepharose beads (Pharmacia) were added. The cells were incubated for 7 days after which the S-Sepharose beads were removed and washed with 0.1 M NaCl in 10 mM Tris buffer (pH 7.5). BPI was eluted from the beads by addition of 1.0 M NaCl in Tris buffer. The top producers, designated Clones 37 and 91, secreted ~ 17 and 14 μ g/ml, respectively in this assay and were frozen as Research Cell Bank numbers C2020 and C2021, respectively. Purified protein was prepared for crystallization studies as follows.

Production and Purification of Nonglycosylated rBPI.

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The host cells used to prepare protein for crystallization studies were CHO-K1 cells transformed with the DNA vector pING4322 which includes DNA encoding the 456 amino acids of human BPI preceded by its endogenous 31 residue secretory signal as described above. During post-translational secretory processing, the signal sequence residues were removed by the host cell. The desired expression product, nonglycosylated rBPI, was a biologically active variant of the human BPI molecule in which the amino acid serine at position 351 in the human BPI protein has been replaced with an alanine.

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Forty roller bottles were prepared which contained the transfected CHO-K1 host cells at 1.3 x 10⁷ cells per bottle in DME/F12 media supplemented with 5% fetal bovine serum (FBS). The cells were grown for three days, at which time 500 ml of fresh media, DME/F12 with 2.5% FBS was added along with a 10 ml slurry (approximately 8 gr.) of sterilized S-Sepharose (Pharmacia, fast flow #17-0511-01, Uppsula, Sweden) and 1 ml of a 1 M solution of sodium butyrate. After two days, the old media plus the S-Sepharose was removed and fresh media, S-Sepharose and sodium butyrate were added to each roller bottle. This process of harvesting the expressed protein product with S-Sepharose was repeated for a total of three harvests, and the S-Sepharose removed during each harvest was pooled. The use of S-Sepharose beads to capture recombinant BPI protein products has been described in U.S. Patent No. 5,439,807 and WO93/23540 (PCT/US93/04752).

The expressed nonglycosylated rBPI protein was purified from the pooled S-Sepharose by first removing it from the S-Sepharose resin followed by further purification and concentration on a series of Q-Sepharose (Pharmacia, fast flow #17-0510-01) and CM-Spherodex (Sepracor, #273431, Villeneuve la Garenne, France) columns. Following purification, the protein was buffer exchanged utilizing a Sephacryl S-100 (Pharmacia, high resolution #17-0612-01) column.

Specifically, the pooled S-Sepharose resin from each harvest was allowed to settle for approximately 15 minutes. The media was removed by decanting and the settled resin was washed three times with approximately 400 ml of 20 mM MES, pH 6.8, 150 mM NaCl. For each wash, the buffer was added, the mixture was stirred gently and the resin was allowed to resettle for approximately 15 minutes. Each buffer wash was removed by decanting. The beads were then washed with 400 mL

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of 20 mM sodium acetate/acetic acid, 150 mM NaCl, pH 4.0 (acetate buffer), and then poured into a 2.5 x 50 cm liquid chromatography column (BIORAD, Econocolumn, Richmond, California). The column was washed extensively with approximately 2 liters of 400 mM NaCl-acetate buffer, pH 4.0 until the A280 absorbance reading of the column eluate was equal to that of the buffer alone. The column was additionally washed with approximately 600 ml of 600 mM NaCl-acetate buffer until the A280 absorbance of the eluate was again equal to that of buffer alone. The protein was then eluted from the S-Sepharose in approximately 500 ml of 1.0 M NaCl-acetate buffer.

The S-Sepharose eluates from each harvest were pooled and diluted to a NaCl concentration of 300 mM. The diluted material was then loaded on to a two column, serial arrangement of a 100 ml Q-Sepharose column connected to a 12 ml CM-Spherodex column. Both columns were constructed using new, sterile resin and were pre-equilibrated with 20 mM MES, pH 5.5, 200 mM NaCl. The Q-Sepharose column served to remove any nucleic acid in the sample material, and the protein did not bind to this resin. After the approximate 3 liters of protein containing material had been loaded, the Q-Sepharose column was disconnected and the CM-Spherodex column was washed with buffer until the A280 absorbance of the eluate was the same as buffer alone. The protein was eluted from the column in 20 mM MES, 400 mM NaCl. pH 5.5 in a volume of approximately 180 ml. This eluted fraction was then reloaded on to a smaller (2 ml) CM-Spherodex column for protein concentration, and the bound protein removed in a single step of 20 mM MES, 1.2 M NaCl, pH 5.5 in a volume of approximately 12 mL. The protein was then loaded directly on to a 150 ml pyrogen-free Sephacryl S-100 column pre-equilibrated with 20 mM sodium citrate, 150 mM NaCl, pH 5.0 buffer. Column fractions were analyzed by Coommassiestained (0.5% Coommassie Brilliant Blue-R, 25% isopropanol, 10% methanol, 10% acetic acid) SDS-PAGE and Western analysis. Western analysis was performed using a 1:1000 dilution of a rabbit anti-human BPI antisera. Fractions which contained the nonglycosylated rBPI protein were pooled and resulted in a final lot which was greater than 95% pure as analyzed by Coommassie-stained SDS-PAGE.

The protein samples thus prepared and purified were filtered and/or concentrated for crystallization studies of the purified nonglycosylated rBPI protein.

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Protein samples were optionally filtered using a 0.2 µm syringe filter (Millipore Corp., Bedford, MA) or a 0.2 µm Nalgene filter (Nalge Corp., Rochester, NY) to remove precipitate. Protein samples were concentrated in a Centricon 10 (Amicon Corp., Beverly, MA) or a Centriprep 10 (Amicon Corp., Beverly, MA). For the Centricon 10 concentrators, a JA 20 rotor (Beckman, Fullerton, CA) in a J2-21 Beckman centrifuge was used at 6000 rpm for 60 minutes. For the Centriprep 10 concentrators, a swinging bucket rotor in a J-6B Beckman centrifuge was used at 3000 rpm for 60 minutes. Final volumes for various protein samples prepared for crystallization studies described herein ranged from about 0.1 to 1 mL, and the protein concentrations were generally between about 10 and 20 mg/mL. Protein solutions may be diluted or concentrated for crystallization studies, for example, from about 5 to about 50 mg/ml.

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Example 2: Structure Determination of a Crystallized BPI Protein

Presented herein is the crystal structure of BPI and two bound phospholipids at 2.4 Å resolution. Our model provides the first structural information on the LPS-binding and lipid transport protein family and suggests a common mode of lipid binding for its members.

Purified, full-length, non-glycosylated, recombinant human BPI expressed in CHO cells was crystallized by hanging-drop vapor diffusion at room temperature. The protein concentration was 8.5 mg/ml and the crystallization buffer contained 12% (w/v) PEG 8000, 200 mM magnesium acetate, and 100 mM sodium cacodylate, pH 6.8. Two crystal forms with slightly different cell dimensions grew under the same conditions in space group C_2 , with one molecule per asymmetric unit. Form 1 crystals were reproducible and had cell dimensions of a=185.0, b=37.2, c=84.3 Å, and $\beta=101.3$ °. Form 2 crystals appeared rarely and had cell dimensions of a=185.6, b=33.0, c=85.2 Å, and $\beta=101.6$ °

For Table 1, x-ray diffraction data were collected at room temperature with the R-AXIS IIC imaging plate area detector mounted on a Rigaku RU200 rotating anode x-ray generator. Data were processed with DENZO and SCALEPACK [Z. Otwinski, in *Proceedings of CCP4 Study Weekend: Data Collection and*

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Processing, L. Sawyer, N. Isaacs, S. Baileys, Eds. (SERC Daresbury Laboratory, Warrington, UK, 1993), pp. 56]. For form 1 crystals, a native data set to 2.8 Å was collected from a single crystal, which was 92.4% complete overall (84.9% complete with an average $I/\sigma(I)=2.3$ in the outermost resolution shell). A native data set to 2.4 Å for form 2 was collected from two crystals and was 92.7% complete overall. (94.6% complete with an average $I/\sigma(I) = 2.6$ in the outermost shell). Because they could be reliably reproduced, form 1 crystals were used for all heavy atom soaks. The structure was solved by multiple isomorphous replacement (MIR) with anomalous scattering. Heavy atom sites were identified by difference-Patterson and difference-Fourier maps. Phase refinement was performed with [Collaborative Computational Project No. 4, Acta Crystallogr., D50:760 (1994)] producing a mean figure of merit (FOM) of 0.57. The MIR map (Fig. 3) was improved by density modification including solvent flattening, histogram matching, and phase extension using DM [Collaborative Computational Project No. 4, supra]. After a partial model was obtained with FRODO [T. A. Jones, J. Appl. Crystallogr., 11:268 (1978)], phase combination was performed with SIGMAA [Collaborative Computational Project No. 4, supra] (final FOM=0.89). CMNP is chloro-Hg-nitrophenol; DMM is dimethyl mercury; PCMBS is parachloromercury-benzene sulfonate; TELA is triethyl lead acetate.

TABLE 1: X-RAY DIFFRACTION DATA

Item	Resolution (Å)	Data Completeness	Rsym* (%)	Sites (N)	RCullis†	Phasing Power	Mid‡
Native 1	2.8	92.4	8.6				
Native 2	2.4	7.26	7.2				
CMNP§	3.2	84.8	6.1	-	0.99	2.04	0.15
DMM	3.5	72.8	9.8	11	65.0	1.49	0.26
PCMBS	3.1	66.4	9.4	3	77.0	1.27	0.38
HgC1,	3.0	86.5	6.9	1	49.0	2.13	0.18
K,PtC1.	3.2	93.3	8.2	3	0.06	89.0	0.13
K,PtBr,	3.1	94.8	5.8	3	73.0	0.88	0.14
TELA#	3.3	94.0	11.3	2	0.98	0.80	0.15
TELA-HgC1,	3.3	91.4	9.6	3	63.0	1.90	0.18
Xenon	3.4	98.2	18.9	5	87.0	69.0	0.18
K ₃ UO ₂ F ₅	3.0	75.0	8.6	2	65.0	1.40	0.16

*R₅ym = 100 (Σ_h | $|_h < |> |)/(\Sigma h$ |h) where < |> is the mean intensity of all symmetry-related reflections |h.

 $+R_{Culls} = (\Sigma | F_{PH} + / \cdot F_P | - F_{H(calc)})/(\Sigma | F_{PH} + / \cdot F_P |)$ for centric reflections, Phasing power $= \{\Sigma | F_{H(calc)}^{\dagger} | / (\Sigma | F_{PH(calc)}^{\dagger} | F_{PH(calc)}^{\dagger} |^{2}]^{1/2}$.

#MID (mean isomorphous difference) = $\Sigma \mid F_{PH}$ - $F_P \mid /\Sigma \mid F_P$, where F_{PH} is the derivative structure factor and F_P is the native structure factor and the sum is over all reflections common to both data sets.

§CMNP, chloro-Hg-nitrophenol; | DMM, dimethyl mercury; IPCMBS, parachloromercurybenzene sulfonate; #TELA, triethyl lead acetate

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Table 2 relates to model refinement and statistics. The model was refined at 2.8 Å through iterative cycles of simulated-annealing with X-PLOR [A. T. Brünger and A. Krukowski, Acta Crystallogr., A46:585 (1990)] and manual rebuilding. 10% of the data were set aside before refinement began for R_{free} [A. T. Brünger Nature, 355:472 (1992)] calculations. When the model had been refined to an R-factor* of 20.4% (R_{free}=32.6%) with the 2.8 Å data, rigid-body minimization was performed against the 2.4 Å data set (R=29.8% to 3.5 Å after minimization). Additional cycles of simulated annealing, positional refinement, correlated individual temperature factor refinement, and manual rebuilding reduced the R-factor to 22.7% and R_{free}=31.3% (no intensity cutoff). An overall anisotropic temperature factor and bulk solvent correction were applied to the observed reflections when R_{free} showed improvement. The model was confirmed by calculating simulated-annealing omit-maps for every part of the structure. The final model contains all 456 residues of the protein, 48 well-ordered waters, and 2 molecules of phosphatidylcholine. Regions of the backbone with poor electron density include residues 148, 232-236, 258-260, and parts of the loop between residues 281-311. Sidechains with poorly defined density were truncated to alanine. The model was examined by the programs PROCHECK [R. A. Laskowski, M. W. McArthur, D. S. Moss, J. M. Thornton, J. Appl. Crystallogr. 26:283 (1993)], VERIFY [R. Lüthy, J. U. Bowie, D. Eisenberg, Nature, 356:83 (1992)], and ERRAT [C. Colovos and T. Yeates, Protein Sci., 2:1511 (1993)].

TABLE 2: REFINEMENT STATISTICS FOR FORM 2 CRYSTALS

	Data		
	Resolution (Å)	2.4	
	Unique reflections (N)	18,898	
5	Completeness (%)	92.7	
	Atoms in model		
	Protein (non-hydrogen)	3532	
	Phosphatidylcholine	102	
	Water	48	
10	Refinement parameters		
	Resolution range (Å)	50.0-2.4	
	R-factor* (%)	22.7	
,	$R_{free}(\%)$	31.3	
	Avg. atomic B factors		
15	Protein	36.9	
	Lipid N, C	49.4, 51.0†	
	Waters	44.6	
	rms‡ deviation from ideality		
	Bonds (Å)	0.006	
20	Angles (deg)	1.4	
	Dihedrals (deg)	26.0	
	Impropers (deg)	1.2	

^{*}R = 100 (Σ | F_{obs} - F_c |)/(Σ F_{obs}) where F_{obs} and F_c are the observed and calculated structure factors, respectively. †B factors for the lipids bound in the NH₂- and COOH-terminal domains.

25 trms=root mean square

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BPI is a boomerang-shaped molecule with approximate dimensions of 135 by 35 Å (Fig. 1, A and B). It consists of two domains of similar size (NH₂- and COOH-terminal) that are connected by a proline-rich linker of 21 residues (positions 230 to 250). The two domains form three structural units; barrels are found at each end of the protein, and a central β sheet forms an interface between the barrels. The secondary structure and topology of the two domains are similar, giving the protein pseudo-twofold symmetry.

Each barrel (residues 10 to 193 and 260 to 421) contains three common structural elements: a short α helix, a five-stranded antiparallel β sheet, and a long helix (Fig. 2A), in that order. We call these elements helix A, sheet N and helix B in the NH₂-terminal domain, and helix A', sheet C and helix B' in the COOH-terminal domain. Sheets N and C have a series of β bulges that change the direction

of their strands and cause a pronounced curve in the sheets. In each domain, the long helix lies along the concave face of the sheet, with the helical axis at $\sim 60^{\circ}$ to the strands of the β sheet. A single disulfide bond between Cys¹³⁵ and Cys¹⁷⁵ anchors helix B to the final strand of sheet N. Situated between the NH₂- and COOH-terminal barrels is a twisted, seven-stranded antiparallel β sheet composed of four strands from the NH₂-terminal domain and three strands from the COOH-terminal domain. This central sheet forms an interface between the two domains and is thus reminiscent of several dimer interfaces stabilized by hydrogen bonds between strands of a β sheet [M. Leeson, B. Henderson, J. Gillig, J. Schwab and J. Smith, *Structure*, 4:253 (1996); D. Ohlendorf, W. F. Anderson, M. Lewis, C. O. Pabo, B. W. Matthews, *J. Mol. Biol.*, 169:757 (1983); G. N. Reeke, J. W. Becker, G. M. Edelman, *J. Biol. Chem.*, 250:1525 (1975)].

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The structural similarity of the two domains of BPI is shown by the superposition [G. H. Cohen, J. Mol. Biol., 190:593 (1986)] in Fig. 2B; they are related by a rotation of 173° and have a root mean square deviation (rmsd) of 3.0 Å on the basis of superposition of 169 C α pairs. The structure shared by these two domains does not resemble other protein folds; several structural alignment programs [N. N. Alexandrov and D. Fischer, Proteins, 25:354 (1996); D. Fischer, C. J. Tsai, R. Nussinov, Protein Eng., 8:981 (1995); L. Holm, C. Sander, Nucl. Acids Res., 22:3600 (1996)] failed to reveal a significant match to any known folds. Significant differences between the superimposed domains are found in two loop regions containing residues 45 and 96 in the NH₂-terminal domain and residues 280 and 348 in the COOH-terminal domain. These differences may be functionally important because the loops around residues 45 and 96 in the NH₂-terminal domain have been implicated in LPS binding and bactericidal activity (see below). This structural similarity of the two domains was unexpected, not only because of their lack of significant sequence identity (<20%), but also because of their functional differences. The NH₂-terminal domain of BPI is cationic and retains the bactericidal, LPS-binding, and LPS-neutralization activities of the intact protein [A. H. Horwitz, et al., Protein Expr. and Purif., 8:28 (1996); C. E. Ooi, J. Weiss, P. Elsbach, B. Frangione and B. Mannion, J. Biol. Chem., 262:14891-14894 (1987); C. E. Ooi, J. Weiss, M. E. Doerfler and P. Elsbach, J. Exp. Med., 174:649 (1991)]. The COOH-terminal

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domain is essentially neutral and shows limited LPS-neutralization activity [S. L. Abrahamson et al., J. Biol. Chem., 272:2149 (1997)]. However, the structural similarity of the two domains may reflect a previously undetected functional similarity: each domain contains a binding pocket for a phospholipid.

After the amino acid sequence had been traced in the electron density maps, two regions of extended electron density remained that could not be accounted for by protein atoms. This density, found in the interior of both domains, was present in the multiple isomorphous replacement (MIR) maps (Fig. 3) at an intensity similar to that of the protein density, and it became the predominant feature in F_{obs} - F_{calc} maps after sequence fitting (both form 1 and form 2 crystals). Electrospray mass spectrometry of the sample used for crystallization revealed two molecules, with relative molecular masses of 522 and 787, in approximately equal amounts. Tandem mass spectrometric analysis was consistent with the two species being phosphoglycerides containing a phosphatidylcholine head group and either one or two 18-carbon acyl chains with one double bond. Phosphatidylcholine (Fig. 4A) is abundant in eukaryotic cells and is presumably bound by BPI in the cells from which the protein is isolated.

The two lipids are bound in extensive apolar pockets on the concave surface of the boomerang, situated between the NH₂-terminal and COOH-terminal barrels and the central β sheet. In the NH₂-terminal domain, the entrance to the pocket is formed by helices A and B. The back and sides are formed by sheet N and the central sheet. The two acyl chains insert ~15 Å into the interior of the protein and are surrounded by apolar side chains (Fig. 4B). The head group lies at the entrance of the pocket and is exposed to solvent. The pocket in the COOH-terminal domain, which has a slightly larger opening, is formed by the analogous secondary structures. Both basic and acidic side chains found near the entrances of the pockets are available for electrostatic interactions with the zwitterionic head group. When the lipids are removed from the model, the pocket in NH₂-terminal domain has a solvent accessible surface area [M. L. Connolly, Science, 221:709 (1983); M. L. Connolly, J. Am. Chem. Soc., 107:1118 (1985)] of 557 Å², and the pocket in the COOH-terminal domain has an area of 413 Å², for a total of 970 Å². The intensity of the electron

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density for the two acyl chains in both pockets is similar and does not indicate whether the single acyl chain species is found predominantly in either pocket.

The discovery of bound phospholipid in our structure suggests a possible site of interaction between BPI and LPS. As seen in Fig. 4A, phosphatidylcholine and LPS share some structural similarity, including negatively charged phosphate groups and, most notably, acyl chains. Since BPI's function is to bind a lipid, LPS, and since lipid is bound in pockets of BPI, it seems reasonable that the acyl chains of LPS bind in the apolar pockets. The following observations support his hypothesis: i) the acyl chains of lipid A are known to be essential for binding by BPI [H. Gazzano-Santoro et al., Infection and Immunity, 63:2201 (1995)]; ii) the binding pockets of BPI are reminiscent of cavities in other lipid-binding proteins [L. Banaszak et al., Adv. Protein Chem., 45:89 (1994)]; and iii) BPI has a significant sequence similarity to two lipid transfer proteins (see below).

Our proposed site of interaction between BPI and the acyl chains of LPS differs from that suggested by previous work focusing on the NH₂-terminal domain. Fragments containing the NH₂-terminal domain of BPI have been identified with equivalent or greater bactericidal and LPS-binding activities relative to the full-length protein [A. H. Horwitz, et al., Protein Expression and Purification, 8:28 (1996); C. E. Ooi, J. Weiss, M. E. Doerfler and P. Elsbach, J. Exp. Med., 174:649 (1991)]. The activity of one NH₂-terminal fragment was reduced when residues past positions 12 or between positions 169 and 199 were deleted [C. Capodici and J. Weiss, J. Immunol., 156:4789 (1996)]. The structure shows that these deletions affect elements of the barrel (at the beginning of helix A and from the middle to the end of helix B) and could significantly alter its structure. While the barrel seems to be the minimal structural unit with full activity, three smaller regions of this domain retain significant LPS-binding, LPS-neutralization, and bactericidal activity [R. G. Little, D. N. Kelner, E. Lim, D. J. Burke and P. J. Conlon, J. Biol. Chem., 268:1865 (1994)]: residues 17-45 (most of helix A and the first β strand of sheet N), residues 82-108 (a β hairpin [residues 82 to 106 of BPI show limited sequence similarity with residues 32 to 51 of the *limulus* anti-LPS factor (LALF), and have been predicted to form an amphipathic β hairpin similar to that seen in the LALF structure [A. Hoess, S. Watson, G. R. Silber and R. Liddington, EMBO 12:3351 (1993)]. Although this

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region of BPI does form a β hairpin, the strict amphipathic character of the loop seen in LALF is not maintained, and a structural superposition shows that the sequence of BPI must be shifted by one residue relative to the proposed sequence alignment] between strands 3 and 4 of sheet N), and residues 142-169 (a segment preceding and including part of helix B). These three regions include 18 basic residues (and only 4 acidic residues) and form a positively charged tip on the NH₂-terminal domain (on the left of Fig.1) which may make favorable electrostatic interactions with negatively charged groups of LPS. Further studies are necessary to determine the relative importance of the apolar pockets and positively charged NH₂-terminal tip to BPI's LPS-binding and bactericidal activities.

BPI is the first member of the mammalian LPS-binding and lipid transfer family to have its three-dimensional structure determined. BPI and LBP are related to two lipid transfer proteins, cholesteryl ester transfer protein (CETP) and phospholipid transfer protein (PLTP) [A. Tall, Annu. Rev. Biochem., 64:235 (1995)]. Alignment of the amino acid sequences of human BPI, LBP, CETP, and PLTP with BPI's secondary structure (Fig. 5, A and B) shows that structurally important residues are conserved in the four proteins. The two cysteines that form the single disulfide bond and are critical to the function of BPI [A. H. Horwitz, et al., Protein Expression and Purification, 8:28 (1996)] are completely conserved. Also, the pattern of hydrophobic/hydrophilic residues in the β strands indicates that the β bulges responsible for the extensive sheet twisting are preserved. The conserved sequences strongly suggest that members of the LPS-binding and lipid transfer family share BPI's two-domain structure and that the two domains are similar in topology.

It is likely that the lipid transfer proteins will also share the apolar binding pockets found in BPI. Striking parallels are found between our BPI-phosphatidylcholine structure and previous work showing that CETP copurifies with an equimolar amount of phosphatidylcholine [A. Tall, Annu. Rev. Biochem., 64:235 (1995)] and has two distinct binding sites [S. Wang, L. Deng, R. W. Milne and A. R. Tall, J. Biol. Chem., 267:17487 (1992)] -- one for neutral lipids and another for phospholipids. The known ligands of CETP and PLTP (cholesteryl esters, triglycerides, retinyl esters and phospholipids) all contain at least one acyl chain which could bind in apolar pockets similar to those in BPI, suggesting a common

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mode of ligand binding in this family. Sequestration of these hydrophobic chains in interior pockets may be critical to the function of the lipid transfer proteins: transfer of apolar ligands in an aqueous environment. Thus, the structure of BPI illuminates the action of the plasma lipid transfer proteins, as well as offering possibilities for how BPI and LBP interact with LPS.

Example 3: Molecular Modeling of BPI Ligands and Mimetics

We have used the information derived from the X-ray crystal structure of BPI presented herein, along with the teachings of the art, including, for example, WO94/20532 (PCT/US94/02465) to design various BPI-related proteins and peptides. These constructs may be divided into categories as illustrated below, including peptides and proteins, including fragments, analogs and variants of the protein, since they best describe the different ways in which different domains and portions may be assembled to achieve new molecules.

1. Individual Peptide Domains: The overlapping BPI peptide data indicated that the N-terminal domain of BPI contains at least three independent functional domains that have one or more of the biological activities of BPI, including, for example, antibacterial, antifungal, anti-heparin and anti-angiogenic activities. Domain I is a region of amino acid residues from about 17 to about 45; Domain II is a region of amino acid residues from about 65 to about 99; and Domain III is a region of amino acid residues from about 142 to about 169. Hundreds of peptide sequences derived from these domains have been synthesized, including addition, deletion and substitution variants of the domain-derived sequences. Through further refinements, smaller "core" regions within these domains have been identified that still retain high levels of activity; for example, within Domain II residues 90-99 and within Domain III residues 148-161.

These peptides have included linear molecules that may or may not assume a conformation that maximally express activity. From the X-ray structure data, segments of BPI are designed that should preserve the three-dimensional structure of these domains when constructed outside the context of the intact protein. For example, both Domains I and II contain hairpin loop structures that are positioned adjacent to one another in space on the proximal tip of the molecule. In contrast,

although Domain III is a helix + turn and not a loop, extensions from both ends of the domain are positioned near enough to each other to consider linking them together. As a result, peptides can be designed that reflect these structures by replacing selected residues in synthetic or recombinant peptides with cysteines, so as to create disulfide-stablized domain mimetics. Since this approach is based upon the actual structure of BPI, it differs from those of other groups that are based upon putative alignments between BPI and the structure of proteins such as the *Limulus* amoebocyte lysate factor (LALF). As examples of these embodiments, listed below are a series of exemplary peptides that, with cysteines added to the N- and C-terminus could assume structures similar to those seen in the intact protein: from Domain I: residues 36-54; from Domain II: residues 84-109, 85-108, 86-107, 87-106, 88-105, 89-104, or 90-103; and from Domain III: residues 142-164.

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- 2. Peptide Domain Hybrids: It has also been demonstrated that certain peptide domain hybrids, which include repeats of the same sequence from a single domain or inter-domain combination of sequences, have enhanced activities. For example, linking Domain II- and III-derived peptides (such as XMP.29: 85-99::148-161) has enhanced biological activity. Interestingly, these domains in the crystal structure are closely associated in space, and peptide XMP.29 may represent a Domain II::III hybrid that actually shares some structural similarity to the intact protein. Based upon the X-ray structure, a Domain II-III peptide that consists of approximately residues 90-103::146-162 is constructed. Such a peptide may even more closely mimic what is seen in the protein.
- 3. BPI "Tip" Mimetics: As discussed above, portions of all three peptide domains discovered and described in WO94/20532 come together on the proximal tip of the N-terminal fragment. As a result, a BPI "tip" mimetic is designed that essentially "slices" off the most extreme tip of the molecule but preserves the critical domain elements. Such a slice would lack the hydrophobic pockets found in the intact protein, but may exhibit activities beyond those of the individual segments. The following segments represent such a "slice" of the three peptide domains. However, to best preserve the geometry between the domains, it would be desirable to insert "linker" sequences between them so as to ensure proper positioning. By fixing these segments in space, programs such as InsightII (Molecular Simulations.

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Inc.) can identify possible linker sequences by i) searching protein databases for similar structures or ii) de novo designing appropriate linkers. In this regard it may be desirable to utilize residues that are not readily susceptible to proteolysis (Ala, Ser, Gly, etc.), or to utilize amino acids like Pro that impose additional spatial constraints on peptide structure. An exemplary peptide consists of Domain I-II-III-derived elements: 37-54:90-104:144-162.

Similarly, by analogy with the above-described cyclic domain structures, the fact that residues 37 and 162 are positioned near each other in the protein suggests that a cyclic tip mimetic could be created by replacing these residues with cysteines, for example, Cys::38-54:90-104:144-161::Cys.

- 4. Extended N-terminal Domains of BPI: The three dimensional structure of BPI indicates that the molecule, which forms N- and C-terminal domains, can be divided into three structural units as described in Example 2. Two of these units represent the N- and C-terminal "barrels" that are formed by residues 10-193 and 260-421, respectively, whereas the third element is a central β sheet structure that forms the interface between the two barrels. Of interest is the fact that the two bound phospholipids in BPI occupy spaces between the two barrel structures and the central β sheet. As a result, the recombinant BPI protein product rBPI₂₁, which essentially contains residues 1-193 lacks some of the structural components necessary to form a complete hydrophobic pocket. A new molecule is constructed that encodes residues 1 to approximately 260 which would contain most of the residues necessary to form a complete hydrophobic pocket.
- 5. Mutants for Immobilizing rBPI₂₁: One application for BPI protein products is their use as affinity removal ligands for endotoxin in solution. For example, immobilizing a BPI protein on a column or membrane matrix would allow removal of endotoxin from endotoxin-contaminated solutions by simply passing those solutions over the immobilized BPI protein. Some of the cysteine-mutated peptides described above may be useful for this purpose, as well as rBPI₂₃. Alternatively, in order to selectively couple a stable, readily produced protein like rBPI₂₁ to a column or membrane, a cysteine could be added to the N- or C-terminus, thus allowing site-specific conjugation and selective orientation of the binding "tip" away from the solid support. Such a construct is alternatively constructed by adding a short linker

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segment (such as Gly-Gly-Gly-Ser) to the C-terminal residue of the BPI protein product, for example, residue 193 of rBPI(1-193), followed by a cysteine residue. Such a construct would have a high probability of folding correctly, given the domain nature of residues 1-193, and be readily conjugatable. Similarly, a series of new conjugates between rBPI(1-193)C and other thiol-containing proteins or molecules is created for the purposes of evaluating new molecules.

PCT/US98/13007

- 6. New N-terminal Dimeric Molecules: As an extension of the above analysis, a new series of N-terminal dimeric molecules can be constructed that take better advantage of the hydrophobic pockets. For example, by replacing the C-terminal barrel with another copy of the N-terminal barrel, an analog of BPI would be created that contains two functional barrels and possibly two functional hydrophobic pockets. One such dimer could be constructed by replacing residues 260-456 with residues 1-193. Alternatively, other more central locations may be identified within the ß sheet structure where symmetry would dictate additional and even better points for duplication.
- 7. C-terminal Fusion Proteins: The C-terminal domains of LBP and CETP appear to mediate interactions with CD14 and lipoproteins. Similarly, the C-terminal domain of BPI has LPS binding and neutralization activities. As a consequence, the C-terminal barrel of BPI (or other family members) could be fused to barrels or domains of other family members and/or to other proteins to alter/modify/enhance their action.
- 8. Homology modeling of BPI family members: The BPI coordinates have been useful in the generation of molecular models of other members of the BPI protein family. Utilizing CLUSTAL (a multiple sequence alignment program in PCGene) and the Homology module (a structure-based homology modeling program in InsightII on the SGI), molecular models (and the corresponding three-dimensional coordinates files) of lipopolysaccharide binding protein (LBP), cholesteryl ester transfer protein (CETP) and phospholipid transfer protein (PLTP) have been generated. With these files, existing mutants are mapped and new ones designed. Published data [Wang et al., *Biochemistry 30*:3484-3490, (1991)] indicate that insertional alterations in three locations of CETP severely impaired cholesteryl ester transfer activity: residues 48-53, residue 165, and residues 373-379. Since residues

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48-53 and residue 165 of CETP coincide structurally with Domains I and III of BPI respectively, the functional domain structure of BPI extends to the other protein family members. Similarly, by virtue of the symmetry between the N- and C-terminal domains, it is likely that the corresponding residues on the C-terminal tip of BPI are involved in recognizing receptors and/or interacting with lipoproteins.

9. Lipid Pocket Mutants: A detailed compilation of the residues in BPI which form the pockets is described in Table 3 as follows. Column 1 of Table 3 indicates the residue name and number. Column 2 shows checked residues which indicate the residues that show a change in solvent accessible surface area with lipid binding. Column 3 shows checked residues that indicate the residues that have some atom within 4 Å of a lipid atom. If the contact is to the head-group of the lipid, the residues are listed at the end, under ENTRY ONLY. Column 4 indicates conservation in 3 BPI and 4 LBP sequences, e.g., for Ile 68, in 3 of the 7 sequences, the residue is similarly Ile; for the other 4 sequences, the residue is Leu (see also column 5). Column 5 indicates alternative residues which occur in BPI or LBP at that position for the 7 BPI and LBP sequences analyzed. Column 6 indicates residues for mutations to block the pockets, using residues selected to be well-conserved (especially in the N-terminal domain) and relatively small. The suggested mutations are all to large sidechains in order to decrease the size of the pocket by as much as possible.

TABLE 3

N-'	N-TERMINAL POCKET - residues contributing to interior											
Residue ¹	ΔSA ²	Within 4Å ³	Conserva- tion ⁴	Alterna- tives ⁵	Mutations ⁶							
Val 7	1	V	2/7	A,T								
Ile 9	V	√	All		w							
Gly 13	V	√	Ali									
Leu 14	V		All									
Tyr 16	V		All									
Ala 17	V	√	All		F							

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	Ser 18	✓	√	1/7	C,A	
	Gly 21	✓	✓	All		F
	Ala 24	✓	√	4/7	V,T,S	Y,H
	Leu 25	√		All		
5	Ile 68	√	√	3/7	L	
	Leu 76	√	✓	All		
	Phe 78	✓	✓	1/7	V,L	
	Leu 117	√	✓	All		
	Leu 119	✓		All ·		
10	Pro 128	√		5/7	A,S	
	Ile 130	√		1/7	v	
	Val 178	√		2/7	L,I	
	Val 182	✓		All		
	Glu 185	✓	✓	1/7 (allele)	K,D,H	
15	Leu 186	✓	✓	All		W
	Tyr 189	√	✓	All		
	Phe 190	√	✓	2/7	V,L	
	Leu 193	V	√	All		
	Leu 220	√	✓	All		
20	Val 222	√		5/7	M,W	
	Val 254	✓	√	6/7	I	
	Leu 256	√		3/7	F	
	Pro 428	√	√	All		
	Thr 429	√	V	1/7	M,L	
25	Pro 430	√	√	5/7	L	·
	Val 433	V		3/7	I	
	Leu 435	√		All		
	Val 453	√		6/7	I	
	ENTRY ONI	Υ				
30	Gln 20	V .	√.	3/7	E	

	Lys 27	V	✓	3/7	R,S	
	Glu 28	V		6/7	K	
	Arg 31	V		2/7	K,E	
	Ser 181	V	V	5/7	T,A	
5	Arg 432	. 1	V	3/7	K,Y,H	
	Tyr 455	√	V	6/7	Н	
,	C-7	ΓERMINA	L POCKE	T - residues	contributing to i	nterior
	Phe 263	V	1	All		
	Asn 264	V		All		
10	Ala 266	V	V	All		F
	Gly 267	V	V	2/7	A,S,T,	
	Val 275	V	1	1/7	A,Y	·
	Leu 276	V	V	5/7	F,W	F,W
	Lys 277	V		1/7	G,N	
15	Met 278	V	V	1/7	L,F	
	Val 318	V		1/7	L,I,G	
	Ala 320	√		2/7	V	
	Pro 324	V	√	6/7	Q	
	Leu 326	V	V	6/7	ν	
20	Phe 335	✓	V	1/7	L,V,E	
	Pro 337	V	V	5/7	A,F	
	Val 339	V		2/7	L,M	
	Met 360	V	V	2/7	L,V	
	Thr 362	✓	V	5/7	L	
25	Val 368	✓ .	V	2/7	I,L	
	Leu 375	√	V	3/7	I,V	
	Val 376	√	V	2/7	I,T	
	Gly 377	V		All		
	Leu 379	√	V	All		
30	Leu 381	V	V	3/7	P	

Val 409	V	V	1/7	L,M,I	
Val 413	V	√	1/7	F,L	F
Val 417	V	√	3/7	I,F	w
Lys 420	V	✓	5/7	Е	Y,H
Leu 421	V	√	5/7	I,F	
Phe 425	V	✓	6/7	L	
ENTRY O	NLY				
Asp 200	✓		All		
Ser 201	V	√	4/7	K,T,N	
Val 202	V		4/7	F,I	
Tyr 270	V	✓	All		
Arg 416	V	V	1/7	K,V,D	
Lys 423	V		3/7	R,E,Q	

- Residue name and number.
- 15 ² Checked residues show a change in solvent accessible surface area with lipid binding.
 - Checked residues have some atom within 4 Å of a lipid atom (if the contact is to the head group of the lipid, the residues are listed at the end, under ENTRY ONLY).
- Conservation in 3 BPI and 4 LBP sequences, e.g., for Ile 68, in 3 of the 7 sequences, the residue is similarly Ile; for the other 4 sequences, the residue is Leu (see note 5).
 - Indicates alternative residues which occur in BPI or LBP at that position for the 7 BPI and LBP sequences analyzed.
- Indicates residues for mutations to block the pockets using residues selected to be well-conserved (especially in the N-terminal domain) and relatively small. The suggested mutations are all to large sidechains in order to decrease the size of the pocket by as much as possible.

							TABLE 4	•		
	ATOM	1	СВ	VAL	A	1	95.942	11.564	18.390	1.00 51.23
-	ATOM	2	CG1	VAL	A	1	97.217	12.313	18.805	1.00 48.90
5	ATOM	3	CG2	VAL	A	1	94.694	12.383	18.720	1.00 50.67
	ATOM	4	С	VAL	A	1	96.124	12.502	16.083	1.00 47.46
10	ATOM	5	0	VAL	A	1	97.238	12.990	15.911	1.00 49.54
	ATOM	8	N	VAL	Α.	1	97.161	10.372	16.554	1.00 51.03
1.5	MOTA	10	CA	VAL	A	1	95.982	11.216	16.876	1.00 48.79
15	ATOM	11	N	ASN	A	2	95.020	13.016	15.554	1.00 42.80
	ATOM	13	CA	ASN	A	2	95.067	14.266	14.810	1.00 38.18
20	MOTA	14	CB	ASN	A	2	93.709	14.566	14.185	1.00 39.01
	ATOM	15	CG	ASN	A	2	93.494	13.831	12.892	1.00 40.06
25	ATOM	16	OD1	ASN	A	2	94.323	13.020	12.487	1.00 42.94
25	ATOM	17	ND2	ASN	A	2	92.380	14.108	12.228	1.00 39.49
	ATOM	20	С	ASN	A	2	95.439	15.376	15.786	1.00 34.97
30	ATOM	21	0	ASN	A	2	94.946	15.399	16.916	1.00 37.36
	ATOM	22	N	PRO	A	3	96.373	16.260	15.397	1.00 30.80
35	ATOM	23	CD	PRO	A	3	97.144	16.244	14.143	1.00 28.90
33	MOTA	24	CA	PRO	A	3	96.806	17.367	16.252	1.00 28.19
	MOTA	25	CB	PRO	A	3	98.083	17.830	15.571	1.00 26.30
40	MOTA	26	CG 	PRO	A	3	97.765	17.619	14.128	1.00 26.17
	MOTA	27	С	PRO	A	3	95.765	18.482	16.283	1.00 28.00
45	ATOM	28.	0	PRO	A	3	95.104	18.758	15.275	1.00 28.89
43	MOTA	29	N	GLY	A .	4	95.615	19.108	17.444	1.00 26.31
	MOTA	31	CA	GLY	A	4	94.653	20.182	17.588	1.00 25.44
50	ATOM '	32	С	GLY	A	4	95.178	21.508	17.091	1.00 25.34
	MOTA	33	0	GLY	A	4	94.407	22.425	16.831	1.00 28.37
55	ATOM	34	N	VAL	A	5	96.494	21.621	16.981	1.00 23.98
	MOTA	36	CA	VAL	À	5	97.134	22.842	16.519	1.00 21.80
	MOTA	37	CB	VAL	A	5	97.671	23.681	17.689	1.00 18.63
60	ATOM	38	CG1	VAL	A	5	98.573	24.785	17.171	1.00 20.04
	ATOM	39	CG2	VAL	A	5	96.526	24.283	18.468	1.00 20.89
65	ATOM	40	С	VAL	A	5	98.308	22.436	15.660	1.00 24.04
05	ATOM	41	0	VAL	A	5	99.014	21.486	15.984	1.00 27.65
	MOTA	42	Ŋ	VAL	A	6	98.492	23.139	14.551	1.00 25.94

	MOTA	44	CA	VAL	A	6	99.589	22.874	13.633	1.00	26.54
	ATOM	45	СВ	VAL .	A	6	99.082	22.292	12.294	1.00	29.26
5	ATOM	46	CG1	VAL	A	6	100.253	22.038	11.357	1.00	31.93
	MOTA	47	CG2	VAL	A	6	98.318	21.001	12.535	1.00	30.52
10	MOTA	48	С	VAL	A	6	100.289	24.191	13.355	1.00	24.99
10	ATOM	49	0	VAL	A	6	99.638	25.202	13.135	1.00	26.97
	MOTA	50	N	VAL	A	7	101.611	24.183	13.416	1.00	26.91
15	MOTA	52	CA	VAL	A	7	102.404	25.371	13.156	1.00	27.18
	ATOM	53	СВ	VAL	A	7	103.401	25.637	14.298	1.00	27.59
20	ATOM	54	CG1	VAL	A	7	104.343	26.773	13.925	1.00	27.21
20	ATOM	5 5	CG2	VAL	A	7	102.659	25.967	15.575	1.00	27.05
	ATOM	56	С	VAL	A	7	103.190	25.103	11.889	1.00	29.61
25	MOTA	5 7	0	VAL	A	7	103.863	24.084	11.777	1.00	32.24
	MOTA	58	N	ARG	A	8	103.077	25.994	10.917	1.00	33.04
30	ATOM	60	CA	ARG	A	8	103.801	25.841	9.668	1.00	33.25
30	MOTA	61	CB	ARG	A	8	102.842	25.631	8.494	1.00	35.70
	ATOM	62	CG	ARG	A	8	102.195	24.244	8.485	1.00	41.64
35	ATOM	63	CD	ARG	A	8	101.309	24.004	7.259	1.00	44.24
	ATOM	64	NE	ARG	A	8	102.049	23.578	6.073	1.00	48.00
40	ATOM	66	CZ	ARG	A	8	101.565	23.626	4.835	1.00	51.86
40	MOTA	67	NH1	ARG	A	8	100.341	24.091	4.611	1.00	52.72
	MOTA	70	NH2	ARG	A	8	102.293	23.177	3.820	1.00	53.10
45	MOTA	73	С	ARG	A	8	104.682	27.052	9.441	1.00	32.06
	MOTA	74	0	ARG	A	8	104.196	28.168	9.288	1.00	32.62
50	MOTA	75	N	ILE	A	9	105.986	26.830	9.515	1.00	31.21
	MOTA	77	CA	ILE	A	9	106.955	27.888	9.315	1.00	31.97
	MOTA	78	СВ	ILE	A	9	108.210	27.633	10.167	1.00	32.76
55	ATOM	79	CG2	ILE	A	9	109.208	28.781	10.012	1.00	29.90
	ATOM	80	CG1	ILE	A	9	107.803	27.454	11.633	1.00	32.76
60	ATOM	81	CD1	ILE	A	9	108.945	27.087	12.554	1.00	36.13
	MOTA	82	С	ILE	A	9	107.309	27.889	7.832	1.00	34.24
	MOTA	83	0	ILE	A	9	107.650	26.845	7.265	1.00	37.01
65	ATOM	84	N	SER	A	10	107.179	29.045	7.195	1.00	34.50

	ATOM	86	CA	SER	A	10	107.473	29.173	5.774	1.00	33.98
	MOTA	87	СВ	SER	A	10	106.536	30.200	5.146	1.00	33.86
5	ATOM	88	OG	SER	A	10	106.778	31.488	5.679	1.00	36.26
	ATOM	90	С	SER	A	10	108.922	29.563	5.500	1.00	32.91
10	ATOM	91	0	SER	A	10	109.674	29.886	6.416	1.00	33.57
10	ATOM	92	N	GLN	A	11	109.300	29.550	4.225	1.00	34.16
	ATOM	94	CA	GLN	A	11	110.653	29.912	3.811	1.00	35.54
15	ATOM	95	СВ	GLN	A	11	110.792	29.845	2.283	1.00	35.39
	MOTA	96	.cg	GLN	A	11	112.212	30.070	1.746	1.00	36.23
20	MOTA	97	CD	GLN	À	11	113.164	28.920	2.040	1.00	35.94
20	ATOM	98	OE1	GLN	A	11	112.880	27.774	1.716	1.00	38.97
	MOTA	99	NE2	GLN	A	11	114.310	29.230	2.624	1.00	35.28
25	ATOM	102	С	GLN	A	11	110.991	31.312	4.316	1.00	36.60
	MOTA	103	0	GLN	A	11	112.116	31.559	4.740	1.00	41.10
30	ATOM	104	N	LYS	A	12	110.013	32.216	4.305	1.00	35.51
30	MOTA	106	CA	LYS	A	12	110.245	33.573	4.788	1.00	31.96
	MOTA	107	CB	LYS	A	12	109.005	34.445	4.612	1.00	32.24
35	MOTA	108	CG	LYS	A	12	109.226	35.876	5.065	1.00	32.71
	ATOM	109	CD	LYS	A	12	107.953	36.685	5.063	1.00	31.87
40	ATOM	110	CE	LYS	A	12	108.227	38.085	5.573	1.00	34.67
40	ATOM	111	NZ	LYS	A	12	106.998	38.914	5.627	1.00	37.88
	MOTA	115	С	LYS	A	12	110.612	33.498	6.259	1.00	30.93
45	MOTA	116	0	LYS	A	12	111.482	34.227	6.728	1.00	32.37
	MOTA	117	N	GLY	A	13	109.961	32.586	6.973	1.00	30.47
50	ATOM	119	CA	GLY	A	13	110.231	32.407	8.385	1.00	27.64
	ATOM	120	С	GLY	A	13	111.593	31.791	8.610	1.00	29.49
	ATOM	121	0	GLY	A	13	112.312	32.184	9.531	1.00	30.35
55	ATOM	122	N	LEU	A	14	111.952	30.817	7.779	1.00	28.87
	ATOM	124	CA	LEU	A	14	113.248	30.170	7.912	1.00	28.53
60	ATOM	125	CB	LEU	A	14	113.361	28.941	7.011	1.00	25.60
	ATOM	126	CG	LEU	A	14	112.564	27.698	7.419	1.00	25.99
	ATOM	127	CD1	LEU	A	14	112.959	26.547	6.531	1.00	28.34
65	MOTA	128	CD2	LEU	A	14	112.829	27.330	8.855		
	ATOM	129	C	LEU	A	14	114.359	31.155	7.613	1.00	28.63

	ATOM	130	0	LEU A		14	115.346	31.207	8.335	1.00	31.86
5	ATOM	131	N	ASP A		15	114.176	31.972	6.582	1.00	31.30
3	MOTA	133	CA	ASP A	١.	15	115.176	32.975	6.215	1.00	33.98
	ATOM	134	CB	ASP A	1	15	114.695	33.832	5.033	1.00	36.03
10	ATOM	135	CG	ASP A		15	114.818	33.127	3.681	1.00	36.52
	MOTA	136	ODI	ASP A		15	115.662	32.209	3.515	1.00	37.01
15	MOTA	137	OD2	ASP A	1	15	114.071	33.530	2.765	1.00	36.25
13	ATOM	138	С	ASP A		15	115.473	33.882	7.413	1.00	35.08
	MOTA	139	0	ASP A	1	15	116.626	34.246	7.654	1.00	37.07
20	MOTA	140	N	TYR A	¥.	16	114.433	34.244	8.162	1.00	34.55
	MOTA	142	CA	TYR A	4	16	114.598	35.098	9.337	1.00	34.48
25	MOTA	143	СВ	TYR A	Ā	16	113.239	35.554	9.865	1.00	34.89
23	MOTA	144	CG	TYR A	Ą	16	113.320	36.661	10.891	1.00	35.28
	ATOM	145	CD1	TYR A	1	16	113.859	37.905	10.562	1.00	36.36
30	ATOM	146	CE1	TYR A	4	16	113.921	38.936	11.496	1.00	34.26
	ATOM	147	CD2	TYR A	Ą	16	112.846	36.474	12.182	1.00	33.51
35	ATOM	148	CE2	TYR A	Ą	16	112.903	37.498	13.118	1.00	34.97
33	MOTA	149	cz	TYR A	Ą	16	113.440	38.724	12.767	1.00	34.08
	ATOM	150	ОН	TYR A	4	16	113.491	39.733	13.697	1.00	37.78
40	MOTA	152	C	TYR A	A	16	115.348	34.340	10.427	1.00	35.46
	MOTA	153	0	TYR A	A	16	116.240	34.883	11.074	1.00	36.90
45	ATOM	154	N	ALA A	A	17	114.982	33.081	10.625	1.00	36.20
.5	MOTA	156	CA	ALA A	A	17	115.632	32.250	11.617	1.00	35.77
	MOTA	157	CB	ALA A	A	17	115.025	30.870	11.615	1.00	33.78
50	MOTA	158	С	ALA 2	A	17	117.118	32.180	11.293	1.00	37.85
	MOTA	159	0	ALA I	A	17	117.957	32.330	12.178	1.00	40.31
55	MOTA	160	Ŋ	SER I	A	18	117.447	32.008	10.017	1.00	39.36
	ATOM	162	CA	SER :	A	18	118.846	31.927	9.612	1.00	42.40
	ATOM	163	CB	SER	A	18	118.993	31.399	8.175	1.00	43.78
60	ATOM	164	OG	SER .	A	18	118.220	32.127	7.236	1.00	48.26
	MOTA	166	С	SER .	A	18	119.605	33.238	9.786	1.00	42.53
65	MOTA	167	0	SER .	A	18	120.768	33.228	10.178	1.00	44.52
UJ.	MOTA	168	N	GLN	A	19	118.959	34.367	9.509	1.00	42.42

	MOTA	170	CA	GLN	A	19	119.630	35.652	9.665	1.00	42.67
	MOTA	171	CB	GLN	A	19	118.805	36.806	9.089	1.00	47.13
5	ATOM	172	CG	GLN	A	19	118.810	36.864	7.563	1.00	57.52
	ATOM	173	CD	GLN	A	19	118.457	38.238	7.012	1.00	62.74
10	ATOM	174	OE1	GLN	A	19	117.488	38.866	7.440	1.00	65.88
10	ATOM	175	NE2	GLN	A	19	119.248	38.710	6.051	1.00	64.58
	ATOM	178	С	GLN	A	19	119.943	35.906	11.126	1.00	40.97
15	ATOM	179	0	GLN	A	19	121.030	36.378	11.456	1.00	41.85
	ATOM	180	N	GLN	A	20	119.008	35.565	12.005	1.00	39.51
20	ATOM	182	CA	GLN	A	20	119.226	35.759	13.430	1.00	37.61
20	ATOM	183	СВ	GLN	A	20	117.944	35.507	14.228	1.00	38.26
	ATOM	184	CG	GLN	A	20	116.764	36.405	13.874	1.00	38.33
25	ATOM	185	CD	GLN	A	20	117.057	37.895	14.002	1.00	41.17
	ATOM	186	OE1	GLN	A	20	116.912	38.642	13.040	1.00	43.42
30	ATOM	187	NE2	GLN	A	20	117.441	38.335	15.192	1.00	42.94
30	MOTA	190	С	GLN	A	20	120.325	34.805	13.878	1.00	36.56
	ATOM	191	0	GLN	A	20	121.208	35.176	14.654	1.00	38.07
35	ATOM	192	N	GLY	A	21	120.285	33.587	13.348	1.00	36.41
	MOTA	194	CA	GLY	A	21	121.276	32.585	13.688	1.00	33.84
40	ATOM	195	С	GLY	A	21	122.665	33.023	13.289	1.00	34.61
10	ATOM	196	0	GLY	A	21	123.520	33.199	14.140	1.00	36.02
	MOTA	197	N	THR	A	22	122.879	33.221	11.994	1.00	35.98
45	MOTA	199	CA	THR	A	22	124.174	33.645	11.462	1.00	39.75
	MOTA	200	CB	THR	A	22	124.036	34.109	9.983	1.00	41.34
50	MOTA	201	OG1	THR	A	22	123.400	33.082	9.212	1.00	40.97
50	ATOM	203	CG2	THR	A	22	125.397	34.408	9.377	1.00	40.54
	ATOM	204	С	THR	A	22	124.780	34.785	12.285	1.00	40.46
55	MOTA	205	0	THR	A	22	125.954	34.741	12.667	1.00	41.11
	ATOM	206	N	ALA	A	23	123.959	35.787	12.578	1.00	41.75
60	ATOM	208	CA	ALA	A	23	124.387	36.944	13.351	1.00	41.89
	MOTA	209	CB	ALA	A	23	123.224	37.895	13.567	1.00	40.51
	ATOM	210	С	ALA	A	23	124.919	36.461	14.678	1.00	43.71
65	ATOM	211	0	ALA	A	23	125.992	36.872	15.114	1.00	47.24
	ATOM	212	N	ALA	A	24	124.175	35.554	15.298	1.00	43.71

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	MOTA	214	CA	ALA	A	24	124.570	34.991	16.574	1.00 43.02
5	ATOM	215	СВ	ALA	A	24	123.413	34.238	17.185	1.00 43.37
3	ATOM	216	С	ALA	A	24	125.774	34.069	16.396	1.00 42.93
	ATOM	217	0	ALA	A	24	126.549	33.868	17.328	1.00 45.41
10	ATOM	218	N	LEU	A	25	125.944	33.532	15.193	1.00 41.61
	ATOM	220	CA	LEU	A	25	127.056	32.638	14.910	1.00 40.32
15	MOTA	221	СВ	LEU	A	25	126.746	31.755	13.699	1.00 36.89
13	ATOM	222	CG	LEU	A	25	127.662	30.554	13.483	1.00 34.95
	MOTA	223	CD1	LEU	A	25	127.636	29.653	14.703	1.00 34.64
20	ATOM	224	CD2	LEU	A	25	127.218	29.797	12.262	1.00 35.16
	MOTA	225	С	LEU	A	25	128.331	33.438	14.676	1.00 40.69
25	MOTA	226	0	LEU	A	25	129.423	32.984	15.014	1.00 42.61
23	MOTA	227	N	GLN	A	26	128.194	34.644	14.138	1.00 41.51
	MOTA	229	CA	GLN	A	26	129.352	35.494	13.878	1.00 45.29
30	MOTA	230	CB	GLN	A	26	128.920	36.850	13.317	1.00 46.09
	MOTA	231	CG	GLN	A	26	130.087	37.811	13.108	1.00 48.70
35	ATOM	232	CD	GLN	A	26	129.650	39.240	12.861	1.00 50.15
<i>J</i> .J	ATOM	233	OEl	GLN	A	26	129.833	40.115	13.708	1.00 51.16
	ATOM	234	NE2	GLN	A	26	129.078	39.489	11.692	1.00 51.55
40	MOTA	237	С	GLN	Α	26	130.175	35.728	15.145	1.00 46.79
	MOTA	238	0	GLN	Α	26·	131.403	35.587	15.136	1.00 46.74
45	MOTA	239	N	LYS	Α	27	129.488	36.081	16.230	1.00 48.40
	MOTA	241	CA	LYS	Α	27	130.140	36.355	17.509	1.00 50.23
	MOTA	242	CB	LYS	A	27	129.119	36.696	18.595	1.00 52.58
50	MOTA	243	ĊG	LYS	A	27	128.157	37.818	18.249	1.00 55.93
	MOTA	244	CD	LYS	A	27	127.495	38.391	19.503	1.00 59.98
55	ATOM	245	CE	LYS	A	27	126.959	37.306	20.437	1.00 63.01
	ATOM	246	NZ	LYS	A	27	125.911	36.452	19.807	1.00 66.76
	MOTA	250	С	LYS	A	27	130.983	35.186	17.977	1.00 50.39
60	MOTA	251	0	LYS	5 A	27 .	132.121	35.369	18.398	1.00 52.00
	MOTA	252	N	GLU	JA	28	130.421	33.984	17.920	1.00 50.09
65	MOTA	254	CA	GLU	JA	28	131.157	32.803	18.341	1.00 48.77
33	MOTA	255	CB	GL	A U	28	130.250	31.571	18.390	1.00 48.51

	MOTA	256	CG	GLU A	A	28	129.514	31.395	19.713	1.00	50.95
	ATOM	257	CD	GLU A	A	28	130.436	30.999	20.857	1.00	52.82
5	MOTA	258	OE1	GLU A	A	28	130.687	29.789	21.030	1.00	55.51
	MOTA	259	OE2	GLU A	A	28	130.903	31.891	21.594	1.00	54.60
10	MOTA	260	С	GLU A	A	28	132.343	32.564	17.421	1.00	46.96
10	ATOM	261	0	GLU A	A	28	133.407	32.153	17.871	1.00	49.87
	ATOM	262	N	LEU A	A	29	132.177	32.868	16.140	1.00	43.03
15	ATOM	264	CA	LEU A	A	29	133.257	32.674	15.189	1.00	39.75
	ATOM	265	CB	LEU A	A	29	132.740	32.817	13.763	1.00	36.79
20	ATOM .	266	CG	LEU A	A	29	131.832	31.658	13.360	1.00	33.73
20	ATOM	267	CD1	LEU A	A	29	131.557	31.708	11.870	1.00	34.40
	ATOM	268	CD2	LEU A	Ą	29	132.499	30.347	13.720	1.00	31.01
25	ATOM	269	С	LEU /	A	29	134.431	33.610	15.444	1.00	38.40
-	ATOM	270	0	LEU A	A .	29	135.584	33.204	15.348	1.00	38.06
30	ATOM	271	N	LYS A	A	30	134.131	34.843	15.828	1.00	38.44
50	ATOM	273	CA	LYS A	A	30	135.155	35,845	16.111	1.00	40.67
	ATOM	274	СВ	LYS A	Ą	30	134.495	37.225	16.210	1.00	41.05
35	ATOM	275	CG	LYS A	A	30	135.448	38.409	16.270	1.00	43.32
	ATOM	276	CD	LYS A	A	30	134.690	39.733	16.198	1.00	44.53
40	ATOM	277	CE	LYS A	A	30	135.633	40.926	16.343	1.00	46.81
	ATOM	278	NZ	LYS A	A	30	134.912	42.238	16.331	1.00	50.27
	ATOM	282	С	LYS A	A	30	135.955	35.539	17.388	1.00	42.29
45	ATOM	283	0	LYS A	A.	30	136.979	36.166	17.654	1.00	42.94
	MOTA	284	N	ARG A	A	31	135.495	34.565	18.168	1.00	44.02
50	MOTA	286	CA	ARG 2	A	31	136.162	34.190	19.411	1.00	44.45
	ATOM	287	CB	ARG A	A	31	135.139	34.005	20.534	1.00	46.86
	ATOM	288	CG	ARG A	A	31	134.504	35.305	20.983	1.00	52.93
55	ATOM	289	CD	ARG A	A	31	133.479	35.095	22.078	1.00	60.04
	ATOM	290	NE	ARG A	A	31	132.947	36.372	22.554	1.00	67.90
60	MOTA	292	CZ	ARG A	A	31	131.839	36.505	23.279	1.00	71.80
	ATOM	293	NH1	ARG A	A	31	131.128	35.437	23.621	1.00	74.30
	ATOM	296	NH2	ARG I	A	31	131.444	37.710	23.672	1.00	73.67
65	ATOM	299	C	ARG 2	A	31	137.008	32.937	19.276	1.00	44.41
	MOTA	300	0	ARG Z	A	31	137.339	32.293	20.273	1.00	46.21

	ATOM	301	N	ILE	A	32	137.343	32.581	18.041	1.00 44.3	0
5	ATOM	303	CA	ILE	A	32	138.162	31.404	17.783	1.00 42.9	6
3	MOTA	304	CB	ILE	A	32	138.054	30.953	16.308	1.00 39.5	0
	ATOM	305	CG2	ILE	A	32	139.146	29.950	15.973	1.00 36.1	2
10	ATOM	306	CG1	ILE	A	32	136.674	30.351	16.052	1.00 37.1	8
	ATOM	307	CD1	ILE	A	32	136.431	29.975	14.624	1.00 37.2	0
16	ATOM	308	С	ILE	A	32	139.613	31.707	18.103	1.00 44.4	4
15	ATOM	309	0	ILE	A	32	140.109	32.794	17.798	1.00 45.6	0
	MOTA	310	N	LYS	A	33	140.288	30.747	18.725	1.00 45.4	5
20	MOTA	312	CA	LYS	A	33	141.692	30.917	19.060	1.00 46.2	0
	MOTA	313	CB	LYS	A	33	141.958	30.569	20.522	1.00 49.3	3
25	ATOM	314	CG	LYS	A	33	141.336	31.546	21.492	1.00 52.4	1
25	MOTA	315	CD	LYS	A	33	141.910	31.387	22.885	1.00 56.1	5
	MOTA	316	CE	LYS	A	33	143.374	31.789	22.918	1.00 59.6	3
30	MOTA	317	NZ	LYS	A	33	143.891	31.817	24.313	1.00 63.3	1
	MOTA	321	С	LYS	A	33	142.619	30.115	18.153	1.00 44.4	.1
35	ATOM	322	0	LYS	A	33	142.563	28.884	18.102	1.00 42.9	9
33	ATOM	323	N	ILE	A	34	143.442	30.836	17.408	1.00 43.4	9
	MOTA	325	CA	ILE	A	34	144.411	30.231	16.512	1.00 43.4	.7
40	MOTA	326	CB	ILE	A	34	144.452	30.958	15.159	1.00 41.9	13
	MOTA	327	CG2	ILE	A	34	143.322	30.464	14.280	1.00 41.3	19
45	MOTA	328	CG1	ILE	A	34	144.347	32.470	15.363	1.00 42.9	₹7
75	MOTA	329	CD1	ILE	A	34	143.967	33.234	14.108	1.00 44.5	59
	MOTA	330	С	ILE	A	34	145.761	30.288	17.216	1.00 44.8	31
50	MOTA	331	0	ILE	A	34	146.142	31.319	17.778	1.00 46.9	€
	ATOM	332	N	PRO	A	35	146.464	29.151	17.267	1.00 43.	35
55	ATOM	333	CD	PRO	A	35	146.008	27.862	16.728	1.00 44.0	53
20	ATOM	334	CA	PRO	A	35	147.773	29.003	17.904	1.00 42.	55
	ATOM	335	CB	PRO	A	35	148.109	27.534	17.650	1.00 42.	05
60	ATOM	336	CG	PRO	A	35	146.778	26.887	17.578	1.00 44.	72
	MOTA	337	С	PRO	A	35	148.860	29.899	17.334	1.00 43.	14
65	MOTA	338	0	PRO	Α	35	148.649	30.640	16.376	1.00 44:	26
05	MOTA	339	N	ASP	Α	36	150.026	29.827	17.959	1.00 42.	32

	ATOM	341	CA	ASP	A	36	151.181	30.584	17.535	1.00	41.85
	ATOM	342	СВ	ASP	A	36	151.933	31.125	18.745	1.00	43.04
5	ATOM	343	CG	ASP	A	36	151.155	32.196	19.485	1.00	45.33
	ATOM	344	OD1	ASP	A	36	150.120	32.666	18.959	1.00	49.40
10	ATOM	345	OD2	ASP	A	36 [°]	151.591	32.584	20.588	1.00	46.99
10	MOTA	346	С	ASP	A	36	152.051	29.612	16. 7 76	1.00	42.29
	ATOM	347	0	ASP	A	36	152.175	28.450	17.159	1.00	43.32
15	MOTA	348	N	TYR	A	37	152.637	30.077	15.687	1.00	42.04
	ATOM	350	CA	TYR .	A	37	153.476	29.228	14.877	1.00	41.59
20	ATOM	351	CB	TYR	A	37	152.979	29.243	13.436	1.00	42.04
20	ATOM.	352	CG	TYR .	A	37	151.564	28.743	13.327	1.00	43.15
	ATOM .	353	CD1	TYR .	A	37	151.295	27.378	13.256	1.00	42.32
25	MOTA	354	CE1	TYR .	A	37	149.991	26.904	13.239	1.00	43.09
	MOTA	355	CD2	TYR	A	37	150.487	29.626	13.370	1.00	44.59
30	ATOM	356	CE2	TYR .	A	37	149.175	29.161	13.354	1.00	43.64
50	ATOM	35 7	CZ	TYR .	A	37	148.938	27.799	13.291	1.00	42.12
	ATOM	358	OH	TYR .	A	37	147.651	27.327	13.307	1.00	43.31
35	MOTA	360	С	TYR	A	37	154.923	29.649	14.957	1.00	42.22
	MOTA	361	0	TYR	A	37	155.311	30.692	14.443	1.00	43.15
40	MOTA	362	N	SER	A	38	155.714	28.833	15.634	1.00	44.23
	MOTA	364	CA	SER	A	38	157.130	29.088	15.789	1.00	47.19
	ATOM	365	CB	SER	A	38	157.466	29.330	17.257	1.00	47.44
45	ATOM	366	OG	SER	A	38	156.651	30.356	17.799	1.00	54.54
	ATOM	368	С	SER	A	38	157.887	27.878	15.285	1.00	48.36
50	ATOM	369	0	SER	A	38	157.664	26.761	15.746	1.00	49.57
	ATOM	370	N	ASP	A	39	158.773	28.103	14.327	1.00	50.18
	ATOM	372	CA	ASP	A	39	159.569	27.030	13.763	1.00	52.06
55	ATOM	373	CB	ASP	A	39	158.822	26.402	12.579	1.00	53.01
	ATOM	374	CG	ASP	A	39	159.375	25.043	12.179	1.00	54.91
60	ATOM	375	OD1	ASP	A	39	159.569	24.178	13.063	1.00	54.67
	ATOM	376	OD2	ASP	A	39	159.600	24.837	10.969	1.00	56.72
	ATOM	3 <i>77</i>	С	ASP	Α	39	160.912	27.631	13.334	1.00	53.93
65	ATOM	378	0	ASP	A	39	161.100	28.853	13.384	1.00	54.67
	ATOM	379	N	SER	Α	40	161.868	26.771	13.003	1.00	54.19

	ATOM	381	CA	SER	A	40	163.187	27.208	12.577	1.00 52.9	4
_	ATOM	382	CB	SER	A	40 .	164.255	26.307	13.202	1.00 52.9	7
5	MOTA	383	OG	SER	A	40	163.996	24.934	12.947	1.00 53.2	7
	MOTA	385	С	SER	A	40	163.283	27.163	11.057	1.00 54.3	2
10	ATOM	386	0	SER	A	40	162.394	26.623	10.386	1.00 54.8	3
	ATOM	387	N	PHE	A	41	164.358	27.730	10.516	1.00 55.1	2
1.6	MOTA	389	CA	PHE	A	41	164.580	27.742	9.074	1.00 55.0	4
15	MOTA	390	СВ	PHE	A	41	163.908	28.967	8.424	1.00 50.9	1
	MOTA	391	CG	PHE	A	41	164.564	30.285	8.756	1.00 46.2	3
20	ATOM	392	CD1	PHE	A	41	165.720	30.695	8.092	1.00 44.9	4
	MOTA	393	CD2	PHE	A	41	164.019	31.124	9.719	1.00 44.3	7
25	ATOM	394	CE1	PHE	A	41	166.321	31.918	8.386	1.00 43.8	5
25	ATOM	395	CE2	PHE	A	41	164.614	32.350	10.018	1.00 43.5	1
	ATOM	396	CZ	PHE	A	41	165.765	32.746	9.351	1.00 42.5	7
30	ATOM	397	С	PHE	A	41	166.071	27.724	8.756	1.00 56.7	5
	ATOM	398	0	PHE	A	41	166.904	27.935	9.637	1.00 58.2	0
35	MOTA	399	N	LYS	A	42	166.391	27.455	7.496	1.00 59.7	1
33	MOTA	401	CA	LYS	A	42	167.768	27.427	7.010	1.00 63.2	:7
	MOTA	402	CB	LYS	A	42	168.474	26.129	7.425	1.00 64.9	8
40	MOTA	403	CG	LYS	A	42	169.984	26.140	7.215	1.00 68.4	6
	ATOM	404	CD	LYS	A	42	170.706	25.591	8.449	1.00 70.9)4
45	ATOM	405	CE	LYS	A	42	172.227	25.630	8.298	1.00 71.6	56
43	MOTA	406	NZ	LYS	A	42	172.928	25.300	9.582	1.00 71.3	L1
	MOTA	410	С	LYS	A	42	167.706	27.552	5.487	1.00 64.2	24
50	MOTA	411	0	LYS	A	42	167.585	26.555	4.773	1.00 65.	07
	MOTA	412	N	ILE	A	43	167.725	28.792	5.003	1.00 64.	98
55	MOTA	414	CA	ILE	A	43	167.652	29.070	3.570	1.00 64.	79
33	MOTA	415	CB	ILE	E A	43	166.721	30.282	3.282	1.00 65.	80
	ATOM	416	CG	2 ILE	E A	43	166.731	30.617	1.796	1.00 66.	80
60	ATOM	417	CG:	1 ILE	E A	43	165.293	29.965	3.745	1.00 64.	11
	MOTA	418	CD:	l ILI	E A	43	164.284	31.071	3.478	1.00 65.	50
65	MOTA	419	С	IL	e a	43	169.039	29.291	2.956	1.00 64.	78
U.S	MOTA	420	0	IL	E A	43	169.840	30.079	3.465	1.00 65.	23

	MOTA	421	N	LYS	A	44	169.273	28.631	1.825	1.00	64.51
	MOTA	423	CA	LYS	A	44	170.539	28.672	1.093	1.00	63.65
5	MOTA	424	CB	LYS	A	44	170.328	28.232	-0.360	1.00	63.66
	ATOM	425	CG	LYS	A	44	170.066	26.742	-0.553	1.00	63.81
10	ATOM	426	CD	LYS	A	44	170.064	26.396	-2.035	1.00	65.27
10	MOTA	427	CE	LYS	A	44	169.910	24.905	-2.289	1.00	65.86
	ATOM	428	NZ	LYS	A	44	169.953	24.599	-3.751	1.00	66.87
15	ATOM	432	C	LYS	A	44	171.382	29.947	1.115	1.00	63.16
	ATOM	433	0	LYS	A	44	172.600	29.876	1.274	1.00	65.20
20	ATOM	434	N	HIS	A	45	170.769	31.107	0.929	1.00	61.08
20	ATOM	436	CA	HIS	A	45	171.550	32.339	0.920	1.00	61.08
	ATOM	437	СВ	HIS	A	45	171.454	33.017	-0.447	1.00	64.62
25	ATOM	438	CG	HIS	Α	45	172.039	32.214	-1.565	1.00	68.00
	ATOM	439	CD2	HIS	A	45	171.695	32.126	-2.871	1.00	68.82
30	ATOM	440	ND1	HIS	A	45	173.124	31.380	-1.397	1.00	70.18
50	ATOM	442	CE1	HIS	A	45	173.425	30.815	-2.552	1.00	71.24
	ATOM	443	NE2	HIS	A	45	172.572	31.250	-3.462	1.00	70.61
35	ATOM	445	C	HIS	A	45	171.167	33.337	1.994	1.00	59.87
	MOTA	446	0	HIS	A	45	171.609	34.486	1.960	1.00	60.80
40	ATOM	447	N	LEU	A	46	170.345	32.900	2.940	1.00	57.18
	MOTA	449	CA	LEU	A	46	169.881	33.767	4.012	1.00	53.69
	MOTA	450	CB	LEU	A	46	168.362	33.699	4.103	1.00	53.23
45	ATOM	451	CG	LEU	A	46	167.715	34.609	5.135	1.00	51.73
	MOTA	452	CD1	LEU	A	46	167.712	36.039	4.628	1.00	52.65
50	MOTA	453	CD2	LEU	A	46	166.310	34.126	5.390	1.00	51.01
	ATOM	454	C	LEU	A	46	170.490	33.336	5.330	1.00	52.64
	ATOM	455	0	LEU	A	46	170.908	34.168	6.136	1.00	52.10
55	MOTA	456	N	GLY	A	47	170.513	32.026	5.550	1.00	51.88
	MOTA	458	CA	GLY	A	47	171.081	31.486	6.768	1.00	51.77
60	ATOM	459	С	GLY	A	47	170.107	30.643	7.564	1.00	51.44
	ATOM	460	0	GLY	A	47	169.059	30.235	7.065	1.00	50.21
	MOTA	461	N	LYS	A	48	170.484	30.357	8.802	1.00	52.15
65	ATOM	463	CA	LYS	A	48	169.664	29.572	9.711	1.00	52.12
	ATOM	464	CB	LYS	A	48	170.497	28.479	10.407	1.00	56.73

	MOTA	465	CG	LYS	A	48	171.878	28.907	10.938	1.00 62.50
-	MOTA	466	CD	LYS	A	48	172.953	28.876	9.843	1.00 65.46
5	ATOM	467	CE	LYS	A	48	174.332	29.252	10.380	1.00 66.23
	MOTA	468	NZ	LYS	A	48	175.380	29.178	9.318	1.00 65.64
10	MOTA	472	С	LYS	A	48	169.041	30.510	10.738	1.00 49.63
	MOTA	473	0	LYS	A	48	169.632	31.533	11.101	1.00 48.85
	MOTA	474	N	GLY	A	49	167.846	30.174	11.200	1.00 46.79
15	ATOM	476	CA	GLY	A	49	167.197	31.026	12.169	1.00 47.22
	ATOM	477	С	GLY	A	49	165.918	30.449	12.728	1.00 47.47
20	MOTA	478	0	GLY	A	49	165.594	29.285	12.506	1.00 46.51
	ATOM	479	N	HIS	A	50	165.166	31.292	13.420	1.00 47.82
25	ATOM	481	CA	HIS	A	50	163.922	30.886	14.041	1.00 48.80
25	ATOM	482	СВ	HIS	A	50	164.126	30.793	15.554	1.00 58.27
	MOTA	483	CG	HIS	A	50	163.658	29.505	16.155	1.00 68.69
30	MOTA	484	CD2	HIS	A	50	164.349	28.453	16.657	1.00 72.98
	MOTA	485	ND1	HIS	A	50	162.324	29.196	16.311	1.00 73.69
35	MOTA	487	CE1	HIS	A	50	162.212	28.010	16.885	1.00 76.58
ננ	MOTA	488	NE2	HIS	A	50	163.426	27.539	17.105	1.00 77.46
	MOTA	490	С	HIS	A	50	162.905	31.965	13.742	1.00 45.89
40	MOTA	491	0	HIS	A	50	163.149	33.137	14.019	1.00 41.84
	MOTA	492	N	TYR	A	51	161.785	31.585	13.143	1.00 46.60
45	ATOM	494	CA	TYR	A	51	160.746	32.555	12.824	1.00 48.52
45	ATOM	495	CB.	TYR	A	51	160.308	32.420	11.354	1.00 50.04
	ATOM	496	CG	TYR	A	51	159.360	31.280	11.098	1.00 51.79
50	ATOM	497	CD1	TYR	A	51	157.987	31.449	11.266	1.00 53.21
	MOTA	498	CE1	TYR	A	51	157.111	30.396	11.103	1.00 56.71
55	MOTA	499	CD2	TYR	A	51	159.833	30.021	10.745	1.00 52.56
33	MOTA	500	CE2	TYR	A	51	158.959	28.955	10.573	1.00 56.44
	MOTA	501	CZ	TYR	A	51	157.597	29.149	10.761	1.00 58.01
60	MOTA	502	ОН	TYR	A	51	156.717	28.094	10.649	1.00 60.76
	ATOM	504	С	TYR	. A	51	159.566	32.330	13.763	1.00 47.31
65	MOTA	505	0	TYR	A	51	159.485	31.291	14.424	1.00 47.36
05	ATOM	506	N	SER	A	52	158.648	33.288	13.812	1.00 46.53

	MOTA	508	CA	SER	A	52	157.480	33.163	14.669	1.00	43.69
	ATOM	509	CB	SER	A	52	157.862	33.449	16.123	1.00	43.87
5	MOTA	510	OG	SER	A	52	156.759	33.233	16.986	1.00	42.88
	ATOM	512	С	SER	A	52	156.344	34.083	14.253	1.00	41.60
10	MOTA	513	0	SER	A	52	156.558	35.261	13.981	1.00	42.86
10	MOTA	514	N	PHE	A	53	155.152	33.513	14.132	1.00	39.20
	MOTA	516	CA	PHE	A	53	153.944	34.260	13.796	1.00	36.73
15	ATOM	517	СВ	PHE	A	53	153.310	33.737	12.507	1.00	33.91
	MOTA	518	CG	PHE	A	53	154.107	34.054	11.271	1.00	32.26
20	ATOM	519	CD1	PHE	A	53	154.039	35.311	10.691	1.00	31.58
20	ATOM	520	CD2	PHE	A	53	154.912	33.092	10.682	1.00	30.53
	MOTA	521	CE1	PHE	A	53	154.759	35.603	9.545	1.00	30.85
25	ATOM	522	CE2	PHE	A	53	155.637	33.373	9.533	1.00	30.45
	MOTA	523	CZ	PHE	A	53	155.559	34.631	8.964	1.00	30.49
30	ATOM	524	C ·	PHE	A	53	153.071	33.965	15.006	1.00	35.40
30	ATOM	525	0	PHE	A	53	152.746	32.814	15.276	1.00	34.75
	MOTA	526	N	TYR	A	54	152.720	34.998	15.757	1.00	34.80
35	ATOM	528	CA	TYR	A	54	151.960	34.798	16.974	1.00	37.29
	ATOM	529	CB	TYR	A	54	152.947	34.533	18.103	1.00	39.38
40	MOTA	530	CG	TYR	A	54	153.841	35.721	18.375	1.00	42.01
40	ATOM	531	CD1	TYR	A	54	154.782	36.139	17.437	1.00	42.10
	ATOM	532	CE1	TYR	A	54	155.552	37.270	17.649	1.00	40.85
45	MOTA	533	CD2	TYR	A	54	153.703	36.467	19.541	1.00	43.83
	MOTA	534	CE2	TYR	A	54	154.472	37.600	19.763	1.00	43.33
50	ATOM	535	CZ	TYR	A	54	155.391	37.995	18.811	1.00	42.17
30	ATOM	536	OH	TYR	A	54	156.139	39.126	19.018	1.00	46.33
	ATOM	538	С	TYR	A	54	151.100	35.990	17.362	1.00	38.69
55	ATOM	539	0	TYR	A	54	151.072	37.007	16.672	1.00	41.66
	ATOM	540	Ň	SER	A	55	150.430	35.858	18.505	1.00	40.04
60	MOTA	542	CA	SER	A	55	149.562	36.894	19.056	1.00	40.41
vv	MOTA	543	СВ	SER	A	55	150.407	38.091	19.499	1.00	41.89
	ATOM	544	OG	SER	Α	55	149.708	38.923	20.409	1.00	46.68
65	ATOM	546	С	SER	Α	55	148.489	37.318	18.051	1.00	41.35
	ATOM	547	0	SER	A	55	148.115	38.492	17.973	1.00	40.80

	ATOM	548	N	MET	A	56	147.984	36.346	17.299	1.00	42.39
5	ATOM	550	CA	MET	A	56	146.968	36.606	16.292	1.00	43.64
3	ATOM	551	СВ	MET	A	56	146.996	35.513	15.225	1.00	42.07
	ATOM	552	CG	MET	A	56	148.310	35.414	14.475	1.00	39.09
10	ATOM	553	SD	MET	A	56	148.335	34.056	13.289	1.00	41.60
	ATOM	554	CE	MET	A	56	149.266	32.847	14.209	1.00	36.68
15	ATOM	555	С	MET	A	56	145.580	36.708	16.910	1.00	46.06
.13	ATOM	556	0	MET	A	56	145.222	35.928	17.794	1.00	48.27
	ATOM	557	N	ASP	A	57	144.820	37.700	16.463	1.00	47.20
20	MOTA	559	CA	ASP	A	57	143.466	37.930	16.941	1.00	47.94
	MOTA	560	СВ	ASP	A	57	143.408	39.167	17.848	1.00	52.70
25	ATOM	561	CG	ASP	A	57	143.578	38.831	19.324	1.00	58.73
20	ATOM	562	OD1	ASP	A	57	142.552	38.587	19.999	1.00	62.99
	MOTA	563	OD2	ASP	A	57	144.729	38.835	19.818	1.00	59.77
30	MOTA	564	C	ASP	A	57	142.566	38.151	15.736	1.00	46.86
	ATOM	565	0	ASP	A	57	143.001	38.684	14.712	1.00	44.98
35	MOTA	566	N	ILE	A	58	141.324	37.699	15.843	1.00	45.46
	ATOM	568	CA	ILE	A	58	140.356	37.872	14.773	1.00	45.28
	ATOM .	569	СВ	ILE	A	58	139.232	36.822	14.869	1.00	42.99
40	ATOM	570	CG2	ILE	A	58	138.200	37.052	13.782	1.00	41.93
	ATOM	571	CG1	ILE	A	58	139.825	35.415	14.744	1.00	42.15
45	MOTA	572	CD1	ILE	A	58	138.803	34.306	14.802	1.00	41.09
	MOTA	573	С	ILE	A	58	139.783	39.277	14.936	1.00	47.07
	ATOM	574	0	ILE	A	58	139.271	39.615	15.999	1.00	48.46
50	MOTA	57 5	N	ARG	Α	59	139.945	40.116	13.918	1.00	49.05
	MOTA	577	CA	ARG	A	59	139.441	41.486	13.973	1.00	51.28
55	MOTA	578	CB	ARG		59	140.435	42.459	13.326	1.00	53.21
	MOTA	579	CG	ARG	Α	59	141.760	42.590	14.062	1.00	55.13
	MOTA	580	CD	ARG	Α	59	141.578	43.175	15.453		59.25
60	ATOM	581	NE	ARG		59	142.833	43.178	16.202	1.00	63.13
	ATOM	583	CZ	ARG	-	59	142.981	43.681	17.424		66.00
65	ATOM	584		ARG			141.950	44.231	18.054		68.47
	ATOM	587	NH2	ARC	A	59	144.164	43.631	18.022	1.00	67.15

	MOTA	590	С	ARG	A	59	138.066	41.627	13.322	1.00 52.12
	MOTA	591	0	ARG	A	59	137.256	42.456	13.746	1.00 54.45
5	MOTA	592	N	GLU	A	60	137.823	40.857	12.265	1.00 52.91
	MOTA	594	CA	GLU	A	60	136.538	40.878	11.570	1.00 52.94
10	MOTA	595	СВ	GLU	A	60	136.492	41.944	10.476	1.00 55.28
10	MOTA	596	CG	GLU	Α	60	135.133	41.994	9.776	1.00 60.74
	MOTA	597	CD	GLU	A	60	135.131	42.841	8.523	1.00 64.33
15	MOTA	598	OE1	GLU	A	60	135.450	44.046	8.618	1.00 67.37
	ATOM	599	OE2	GLU	A	60	134.798	42.300	7.443	1.00 65.16
20	_ ATOM	600	С	GLU	A	60	136.279	39.518	10.948	1.00 51.42
20	MOTA	601	0	GLU	A	60	137.170	38.930	10.342	1.00 52.44
	ATOM	602	N	PHE	A	61	135.052	39.035	11.093	1.00 50.66
25	MOTA	604	CA	PHE	A	61	134.646	37.741	10.558	1.00 49.14
	ATOM	605	СВ	PHE	A	61	134.536	36.736	11.707	1.00 44.10
30	ATOM	606	CG	PHE	A	61	134.679	35.308	11.289	1.00 37.41
30	MOTA	607	CD1	PHE	A	61	133.998	34.814	10.188	1.00 36.68
	MOTA	608	CD2	PHE	A	61	135.486	34.448	12.018	1.00 34.73
35	ATOM	609	CEI	PHE	A	61	134.116	33.477	9.817	1.00 36.53
	ATOM	610	CE2	PHE	A	61	135.612	33.115	11.658	1.00 34.86
40	MOTA	611	CZ	PHE	A	61	134.922	32.627	10.553	1.00 35.68
70	ATOM	612	С	PHE	A	61	133.278	37.962	9.924	1.00 50.95
	ATOM	613	0	PHE	A	61	132.249	37.646	10.521	1.00 52.86
45	ATOM	614	N	GLN	A	62	133.259	38.540	8.729	1.00 51.57
	ATOM	616	CA	GLN	A	62	131.993	38.813	8.080	1.00 51.07
50	ATOM	617	CB	GLN	A	62	132.097	39.993	7.120	1.00 54.59
	MOTA	618	CG	GLN	A	62	130.727	40.463	6.639	1.00 62.31
	MOTA	619	CD	GLN	A	62	130.775	41.665	5.708	1.00 67.35
55	MOTA	620	OE1	GLN	A	62	129.812	41.935	4.991	1.00 69.86
	MOTA	621	NE2	GLN	A	62	131.884	42.401	5.727	1.00 71.01
60	MOTA	624	С	GLN	A	62	131.391	37.625	7.367	1.00 48.52
- •	MOTA	625	0	GLN	A	62	132.065	36.912	6.629	1.00 48.13
	MOTA	626	N	LEU	A	63	130.107	37.422	7.622	100 48.11
65	MOTA	628	CA	LEU	Α	63	129.324	36.358	7.017	1.00 47.47
	MOTA	629	CB	LEU	A	63	128.753	35.438	8.103	1.00 44.43

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	MOTA	630	CG	LEU A	A	63	129.754	34.759	9.044	1.00 4	2.13
_	MOTA	631	CD1	LEU A	A.	63	129.021	34.080	10.181	1.00 4	2.20
5	MOTA	632	CD2	LEU 2	A	63	130.593	33.752	8.282	1.00 4	0.96
	MOTA	633	С	LEU 2	A	63	128.208	37.122	6.314	1.00 4	8.35
10	MOTA	634	0	LEU 2	A	63	127.138	37.345	6.882	1.00 4	9.30
	MOTA	635	N	PRO 2	A	64	128.471	37.585	5.082	1.00 4	9.75
1.5	MOTA	636	CD	PRO .	A	64	129.686	37.302	4.297	1.00 5	0.27
15	ATOM	637	CA	PRO .	A	64	127.508	38.347	4.283	1.00 5	0.73
	ATOM	638	СВ	PRO .	A	64	128.309	38.687	3.026	1.00 5	0.48
20	ATOM	639	CG	PRO	A	64	129.210	37.513	2.878	1.00 4	9.49
	ATOM	640	С	PRO	A	64	126.219	37.616	3.937	1.00 5	1.98
25	ATOM	641	0	PRO	A	64	125.129	38.167	4.083	1.00 5	4.28
25	ATOM	642	N	SER	A	65	126.341	36.375	3.488	1.00 5	1.59
	ATOM	644	CA	SER	A	65	125.177	35.607	3.101	1.00 5	0.95
30	ATOM	645	СВ	SER	A	65	125.217	35.356	1.599	1.00 5	2.70
	MOTA	646	OG	SER	A	65	126.503	34.910	1.205	1.00 5	55.07
35	MOTA	648	C	SER	A	65	125.053	34.288	3.832	1.00 5	0.87
J J	ATOM	649	0	SER	A	65	126.052	33.627	4.120	1.00 5	2.05
	MOTA	650	N	SER	A	66	,123.811	33.928	4.136	1.00 5	50.53
40	ATOM	652	CA	SER	A	66	123.477	32.680	4.805	1.00 4	18.85
	ATOM	653	СВ	SER	A	66	123.322	32.878	6.312	1.00 4	18.24
45	ATOM	654	OG	SER	A	66	122.307	33.820	6.615	1.00 4	19.82
1.5	ATOM	656	C	SER	A	66	122.154	32.244	4.202	1.00	49.43
	MOTA	657	0	SER	A	66	121.365	33.080	3.745	1.00	48.59 °
50	MOTA	658	N	GLN	A	67	121.925	30.940	4.165	1.00	49.88
	MOTA	660	CA	GLN	A	67	120.696	30.409	3.605	1.00	49.90
55	MOTA	661	CB	GLN	A	67	120.831	30.267	2.086	1.00	52.61
	ATOM	662	CG	GLN	A	67	121.928	29.308	1.632	1.00	58.79
	ATOM	663	CD	GLN	A	67	121.984	29.154	0.121	1.00	62.57
60	ATOM	664	OE:	1 GLN	A	67	122.550	29.997	-0.576	1.00	66.32
	MOTA	665	NE:	2 GLN	A	67	121.407	28.073	-0.392	1.00	62.89
65	MOTA	668	С	GLN	Α	67	120.337	29.065	4.221	1.00	48.02
J .	ATOM	. 669	0	GLN	Α	67	121.215	28.306	4.635	1.00	49.44

	MOTA	670	N	ILE .	A	68	119.040	28.803	4.328	1.00	45.23
	ATOM	672	CA	ILE .	A	68	118.551	27.544	4.864	1.00	42.63
5	MOTA	673	СВ	ILE .	A	68	117.826	27.721	6.220	1.00	44.01
	MOTA	674	CG2	ILE .	A	68	116.977	26.499	6.546	1.00	45.24
10	ATOM	675	CG1	ILE .	A	68	118.850	27.925	7.333	1.00	44.56
10	MOTA	676	CD1	ILE .	A	68	118.260	27.851	8.719	1.00	47.20
	MOTA	677	С	ILE .	Α	68	117.613	26.935	3.835	1.00	40.01
15	ATOM	678	0	ILE .	A	68	116.560	27.491	3.524	1.00	41.30
	MOTA	679	N	SER .	A	69	118.051	25.833	3.250	1.00	37.17
20	MOTA	681	CA	SER .	A	69	117.264	25.140	2.258	1.00	34.46
20	ATOM	682	СВ	SER	A	69	118.165	24.600	1.147	1.00	34.89
	MOTA	683	OG	SER	A	69	118.890	25.640	0.511	1.00	40.63
25	ATOM	685	С	SER .	A	69	116.570	23.984	2.939	1.00	33.71
	ATOM	686	0	SER	A	69	117.085	23.414	3.896	1.00	34.03
30	ATOM	687	N	MET .	A	70	115.405	23.625	2.430	1.00	33.11
30	MOTA	689	CA	MET	A	70	114.659	22.519	2.981	1.00	34.74
	MOTA	690	СВ	MET	A	70	113.166	22.841	3.012	1.00	36.59
35	MOTA	691	CG	MET .	A	70	112.794	23.880	4.051	1.00	41.38
	MOTA	692	SD	MET	A	70	111.050	24.288	4.028	1.00	47.62
40	MOTA	693	CE	MET	A	70	111.101	25.834	3.173	1.00	47.72
40	MOTA	694	С	MET	A	70	114.906	21.274	2.153	1.00	35.74
	ATOM	695	0	MET	A	70	114.752	21.279	0.928	1.00	37.33
45	ATOM	696	N	VAL	A	71	115.366	20.226	2.820	1.00	35.02
	MOTA	698	CA	VAL	A	71	115.613	18.957	2.165	1.00	32.82
50	ATOM	699	CB	VAL	A	71	116.938	18.343	2.637	1.00	30.47
50	ATOM	700	CG1	VAL	Α	71	117.206	17.049	1.905	1.00	28.06
	ATOM	701	CG2	VAL	A	71	118.069	19.324	2.404	1.00	28.88
55	ATOM	702	С	VAL	A	71	114.438	18.072	2.571	1.00	33.73
	ATOM	703	0	VAL	A	71	114.430	17.505	3.664	1.00	35.31
60	ATOM	704	N	PRO	A	72	113.411	17.975	1.708	1.00	33.98
	ATOM	705	CD	PRO	A	72	113.402	18.500	0.327	1.00	33.44
	MOTA	706	CA	PRO	A	72	112.211	17.168	1.965	1.00	34.72
65	ATOM	707	CB	PRO	A	72	111.660	16.936	0.562	1.00	33.90
	MOTA	708	CG	PRO	A	72	111.980	18.231	-0.124	1.00	32.49

	MOTA	709	С	PRO	A	72	112.495	15.857	2.695	1.00 36.34
	ATOM	710	0	PRO	A	72	113.417	15.131	2.340	1.00 39.70
5	MOTA	711	N	ASN	A	73	111.745	15.607	3.765	1.00 38.08
	ATOM	713	CA	ASN	A	73	111.886	14.399	4.580	1.00 39.44
10	ATOM	714	СВ	ASN	A	73	111.553	13.146	3.768	1.00 45.34
	ATOM	715	CG	ASN	A	73	110.071	12.875	3.699	1.00 50.49
16	ATOM	716	OD1	ASN	A	73	109.392	13.317	2.770	1.00 53.54
15	ATOM	717	ND2	ASN	A	73`	109.552	12.149	4.690	1.00 53.95
	MOTA	720	С	ASN	A	73	113.231	14.192	5.252	1.00 38.00
20	MOTA	721	0	ASN	A	73	113.408	13.204	5.962	1.00 37.89
	MOTA	722	N	VAL	A	74	114.165	15.116	5.050	1.00 37.67
25	ATOM	724	CA	VAL	A	74	115.498	14.995	5.632	1.00 36.66
23	MOTA	725	СВ	VAL	A	74	116.598	15.041	4.525	1.00 37.47
	MOTA	726	CG1	VAL	A	74	117.976	14.753	5.118	1.00 35.10
30	MOTA	727	CG2	VAL	A	74	116.280	14.046	3.402	1.00 33.69
	ATOM	728	С	VAL	A	74	115.784	16.054	6.704	1.00 37.30
35	MOTA	729	0	VAL	A	74	116.106	15.717	7.848	1.00 39.11
33	ATOM	730	N	GLY	A	75	115.672	17.328	6.342	1.00 35.75
	ATOM	732	CA	GLY	A	75	115.928	18.384	7.306	1.00 33.77
40	ATOM	733	С	GLY	A	75 :	116.279	19.698	6.647	1.00 33.37
	ATOM	734	0	GLY	A	75	115.827	19.981	5.536	1.00 32.85
45	ATOM	735	N	LEU	A	76	117.075	20.509	7.338	1.00 34.22
	MOTA	737	CA	LEU	A	76	117.504	21.811	6.828	1.00 33.33
	MOTA	738	CB	LEU	Α	76	117.329	22.894	7.897	1.00 30.43
50	MOTA	739	CG	LEU	Α	76	115.958	23.105	8.524	1.00 28.67
	ATOM	740	CD1	LEU	Α	76	116.081	24.069	9.683	1.00 29.06
55	ATOM	741	CD2	2 LEU	Α	76	114.995	23.633	7.481	1.00 31.34
	ATOM	742	С	LEU	A	76	118.979	21.755	6.448	1.00 33.33
	ATOM	743	0	LEU	J A	76	119.736	20.942	6.967	1.00 34.06
60	ATOM	744	N	LYS	A	77	119.383	22.636	5.550	1.00 34.34
	MOTA	746	CA	LYS	A	77	120.766	22.711	5.131	1.00 33.56
65	MOTA	747	CB	LYS	S A	77	120.927	22.201	3.700	1.00 35.32
	MOTA	748	CG	LYS	S A	77	122.345	22.283	3.166	1.00 38.99

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	MOTA	749	CD	LYS A	77		122.450	21.670	1.782	1.00	41.42
	MOTA	750	CE	LYS A	77	,	122.388	20.150	1.838	1.00	41.64
5	MOTA	751	NZ	LYS A	77	,	123.630	19.565	2.421	1.00	43.94
	ATOM	755	С	LYS A	77	•	121:131	24.175	5.228	1.00	34.11
10	MOTA	. 756	0	LYS A	77	•	120.556	25.013	4.538	1.00	34.01
10	ATOM	.757	N	PHE A	78	}	122.009	24.478	6.173	1.00	36.93
	MOTA	759	CA	PHE A	78	}	122.482	25.830	6.427	1.00	37.23
15	ATOM	760	CB	PHE A	78	3	122.720	25.983	7.933	1.00	37.11
	ATOM	761	CG	PHE A	78	3	123.233	27.329	8.340	1.00	36.19
••	ATOM	762	CD1	PHE A	78	3	122.536	28.481	8.016	1.00	35.10
20	ATOM	763	CD2	PHE A	78	3	124.415	27.442	9.059	1.00	36.97
	ATOM	764	CE1	PHE A	. 78	3	123.008	29.725	8.400	1.00	33.97
25	ATOM	765	CE2	PHE A	. 78	3	124.892	28.679	9.446	1.00	36.53
	ATOM	766	cz	PHE A	. 78	3	124.185	29.824	9.115	1.00	36.23
20	ATOM	767	С	PHE A	. 78	3	123.780	26.063	5.646	1.00	38.21
30	ATOM	768	0	PHE A	. 78	3	124.706	25.256	5.731	1.00	37.35
	ATOM	769	N	SER A	. 79	€	123.845	27.160	4.892	1.00	39.48
35	MOTA	771	CA	SER A	. 79	Ð	125.028	27.489	4.097	1.00	40.20
	ATOM	772	СВ	SER A	. 79	9	124.802	27.126	2.625	1.00	41.51
40	MOTA	773	OG	SER A	. 79	9	i24.408	25.773	2.464	1.00	48.40
40	MOTA	775	С	SER A	. 79	9	125.416	28.963	4.160	1.00	40.10
	ATOM	776	0	SER A	. 79	9	124.557	29.842	4.112	1.00	39.14
45	ATOM	7 77	N	ILE A	8	0	126.713	29.219	4.297	1.00	40.97
	ATOM	779	CA	ILE P	8	0	127.263	30.575	4.309	1.00	41.40
50	ATOM	780	CB	ILE A	8	0	128.002	30.885	5.612	1.00	39.45
50	ATOM	781	CG2	ILE A	8	0	128.518	32.307	5.583	1.00	40.67
	ATOM	782	CG1	ILE A	8	0	127.057	30.714	6.797	1.00	38.41
55	ATOM	783	CD1	ILE A	8	0	127.690	31.022	8.120	1.00	39.45
	ATOM	784	C	ILE A	8	0	128.246	30.524	3.143	1.00	43.93
60	ATOM	785	0	ILE A	8	0 .	128.944	29.519	2.976	1.00	45.50
60	MOTA	786	N	SER I	A 8	1	128.327	31.585	2.344	1.00	47.35
	MOTA	788	CA	SER A	A 8	1	129.193	31.541	1.168	1.00	49.22
65	MOTA	789	СВ	SER I	A _ 8	1	128.338	31.613	-0.097	1.00	49.49
	MOTA	790	OG	SER I	A 8	1	127.332	32.602	0.032	1.00	52.43

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	ATOM	792	С	SER	A	81	130.412	32.432	1.004	1.00	50.65
5	ATOM	793	0	SER	A	81	131.482	31.937	0.658	1.00	52.88
3	ATOM	794	N	ASN	A	82	130.276	33.738	1.185	1.00	52.50
	ATOM	796	CA	ASN	A	82	131.442	34.599	0.995	1.00	54.71
10	ATOM	797	СВ	ASN	A	82	131.104	35.767	0.061	1.00	58.34
	MOTA	798	CG	ASN	A	82	132.342	36.391	-0.570	1.00	61.43
15	ATOM	799	OD1	ASN	A	82	132.462	37.613	-0.657	1.00	63.94
13	MOTA	800	ND2	ASN	A	82	133.257	35.550	-1.039	1.00	61.92
	ATOM	803	С	ASN	A	82	132.040	35.098	2.303	1.00	54.00
20	ATOM	804	0	ASN	A	82	132.268	36.297	2.482	1.00	54.15
	MOTA	805	N	ALA	A	83	132.325	34.162	3.205	1.00	53.37
25	MOTA	807	CA	ALA	A	83	132.892	34.501	4.506	1.00	52.11
23	ATOM	808	СВ	ALA	A	83	132.952	33.273	5.402	1.00	51.81
	ATOM	809	С	ALA	A	83	134.272	35.131	4.359	1.00	50.64
30	MOTA	810	0	ALA	A	83	135.124	34.640	3.610	1.00	49.58
	ATOM	811	N	ASN	A	84	134.467	36.228	5.078	1.00	48.85
35	MOTA	813	CA	ASN	A	84	135.706	36.983	5.063	1.00	47.97
33	MOTA	814	CB	ASN	A	84	135.423	38.393	4.536	1.00	51.32
	MOTA	815	CG	ASN	A	84	136.560	39.361	4.793	1.00	56.05
40	ATOM	816	OD1	asn	A	84	137.470	39.499	3.978	1.00	61.24
	MOTA	817	ND2	asn	A	84	136.500	40.060	5.919	1.00	56.91
45	MOTA	820	С	ASN	A	84	136.243	37.042	6.485	1.00	45.82
	MOTA	821	0	ASN	A	84	135.601	37.606	7.368	1.00	47.29
	ATOM	822	N	ILE	A	85	137.405	36.438	6.703	1.00	43.85
50	MOTA	824	CA	ILE	A	85	138.052	36.402	8.014	1.00	41.41
	MOTA	825	CB	ILE	A	85	138.495	34.952	8.386	1.00	35.61
55	MOTA	826	CG2	ILE	A	85	139.011	34.901	9.804	1.00	32.01
	MOTA	827	CG1	ILE	A	85	137.342	33.965	8.222	1.00	30.72
	MOTA	828	CD1	ILE	A	85	137.712	32.557	8.587	1.00	25.79
60	MOTA	829	С	ILE	Α	85	139.315	37.272	8.017	1.00	43.73
	MOTA	830	0	ILE	A	85	140.343	36.873	7.471	1.00	47.66
65	ATOM	831	N	LYS	A	86	139.239	38.471	8.578	1.00	42.83
	ATOM	833	CA	LYS	A	86	140.420	39.313	8.645	1.00	43.65

	ATOM	834	СВ	LYS A	86	140.090	40.779	8.363	1.00	47.39
	MOTA	835	CG	LYS A	86	139.895	41.055	6.873	1.00	54.31
5	ATOM	836	CD	LYS A	86	140.065	42.528	6.509	1.00	58.05
	ATOM	837	CE	LYS A	86	138.937	43.394	7.045	1.00	61.18
10	ATOM	838	NZ	LYS A	86	139.107	44.820	6.642	1.00	64.32
10	ATOM	842	С	LYS A	86	141.100	39.135	9.996	1.00	43.61
	ATOM	843	0	LYS A	86	140.514	39.404	11.043	1.00	42.95
15	ATOM	844	N	ILE A	87	142.327	38.627	9.961	1.00	43.89
	MOTA	846	CA	ILE A	87	143.104	38.373	11.165	1.00	44.27
20	ATOM	847	СВ	ILE A	87	143.568	36.895	11.206	1.00	43.55
20	ATOM	848	CG2	ILE A	87	144.365	36.609	12.473	1.00	42.71
	ATOM	849	CG1	ILE A	87	142.358	35.964	11.129	1.00	41.65
25	MOTA	850	CD1	ILE A	87	142.723	34.504	11.044	1.00	42.75
	ATOM	851	С	ILE A	87	144.327	39.291	11.212	1.00	44.82
20	MOTA	852	0	ILE A	87	144.817	39.745	10.176	1.00	46.46
30	ATOM	853	N	SER A	88	144.805	39.568	12.417	1.00	43.43
	MOTA	855	CA	SER A	. 88	145.970	40.414	12.606	1.00	44.37
35	MOTA	856	CB	SER A	. 88	145.548	41.825	13.027	1.00	46.30
	MOTA	857	OG	SER A	88	144.757	42.451	12.029	1.00	50.16
40	MOTA	859	С	SER A	. 88	146.830	39.795	13.690	1.00	43.85
40	MOTA	860	0	SER A	. 88	146.327	39.073	14.549	1.00	43.41
	ATOM	861	N	GLY A	89	148.128	40.061	13.638	1.00	45.04
45	MOTA	863	CA	GLY A	. 89	149.029	39.525	14.640	1.00	47.03
	ATOM	864	С	GLY A	. 89	150.415	40.123	14.532	1.00	47.50
50	MOTA	865	0	GLY A	. 89	150.611	41.140	13.861	1.00	47.81
50	MOTA	866	N	LYS A	90	151.377	39.497	15.200	1.00	47.62
	MOTA	868	CA	LYS A	90	152.759	39.957	15.176	1.00	47.17
55	MOTA	869	СВ	LYS A	90	153.197	40.427	16.567	1.00	47.20
	ATOM	870	CG	LYS A	90	152.328	41.514	17.189	1.00	49.57
60	MOTA	871	CD	LYS A	90	152.708	41.723	18.651	1.00	53.99
00	ATOM	872	CE	LYS A	90	151.725	42.625	19.391	1.00	57.45
	ATOM	873	NZ	LYS A	90	152.042	42.735	20.855	1.00	59.91
65	ATOM	877	С	LYS A	90	153.629	38.787	14.738	1.00	45.69
	ATOM	878	0	LYS A	90	153.171	37.644	14.697	1.00	45.87

ATOM TRP A 91 879 N 154.874 39.078 14.389 1.00 44.12 MOTA 881 CA TRP A 91 155.817 38.052 13.969 1.00 43.19 5 ATOM 882 CB TRP A 91 155.734 37.824 1.00 39.90 12.456 MOTA 883 CG TRP A 91 156.078 39.038 11.694 1.00 37.85 10 39.466 MOTA 884 CD2 TRP A 91 157.390 11.313 1.00 39.20 MOTA 885 CE2 TRP A 91 157.251 40.722 10.682 1.00 39.17 MOTA 886 CE3 TRP A 91 158.672 38.918 11.451 1.00 36.54 15 MOTA 887 CD1 TRP A 91 155.222 40.012 11.286 1.00 39.46 ATOM 888 NE1 TRP A 91 155.916 41.032 10.681 1.00 39.30 20 ATOM 890 CZ2 TRP A 91 158.344 41.437 10.188 - 1.00 38.03 CZ3 TRP A 159.757 39.628 10.963 MOTA 891 91 1.00 35.59 MOTA 892 CH2 TRP A 91 159.585 40.876 10.338 1.00 36.22 25 ATOM 893 C TRP A 91 157.218 38.520 14.339 1.00 43.43 MOTA 894 0 TRP A 91 157.460 39.717 14.509 1.00 44.14 30 MOTA LYS A 895 N 92 158.136 37.575 14.468 1.00 44.32 MOTA 897 CA LYS A 92 159.517 37.892 14.789 1.00 44.34 MOTA 898 CB LYS A 92 159.717 38.076 16.302 1.00 47.57 35 MOTA 899 LYS A 92 159.252 36.933 1.00 53.21 CG 17.200 MOTA 900 CD LYS A 92 159.361 1.00 57.24 37.354 18.678 40 MOTA 901 CE LYS A 158.844 1.00 59.23 92 36.283 19.650 **ATOM** 902 ΝZ LYS A 92 158.816 36.742 21.083 1.00 59.30 MOTA 906 C LYS A 92 160.409 36.801 14.227 1.00 42.18 45 ATOM 907 LYS A 92 160.052 35.623 14.255 1.00 43.11 0 MOTA 908 ALA A 93 N 161.521 37.214 13.632 1.00 39.24 50 1.00 36.99 **ATOM** 910 CA ALA A 93 162.471 36.294 13.033 **ATOM** 911 CB ALA A 162.410 36,392 11.515 1.00 36.10 MOTA 912 С ALA A 93 163.850 36.675 1.00 37.81 13.525 55 ATOM 913 0 ALA A 93 164.130 37.851 13.747 1.00 39.61 164.706 **ATOM** 914 N GLN A 94 35.679 13.705 1.00 39.81 60 MOTA 916 CA GLN A 94 166.060 35.919 1.00 41.70 14.173 MOTA 917 CB GLN A 94 166.184 35.541 15.647 1.00 45.63 MOTA 918 CG GLN A 94 167.552 35.803 16.237 1.00 55.83 65 ATOM 919 CD GLN A 94 167.559 35.745 17.750 1.00 63.01

	ATOM	920	0E1	GLN	A	94	168.328	36.456	18.406	1.00	67.41
	ATOM	921	NE2	GLN	A	94	166.703	34.899	18.320	1.00	66.99
5	ATOM	924	С	GLN	A	94	167.067	35.139	13.340	1.00	41.13
	ATOM	925	0	GLN	A	94	166.952	33.921	13.194	1.00	41.48
10	ATOM	926	N	LYS	A	95	168.024	35.865	12.771	1.00	41.51
10	MOTA	928	CA	LYS	A	95	169.084	35.302	11.938	1.00	40.50
	MOTA	929	СВ	LYS	A	95	169.000	35.901	10.531	1.00	40.86
15	MOTA	930	CG	LYS	A	95	170.099	35.478	9.580	1.00	44.00
	ATOM	931	CD	LYS	A	95	169.849	36.030	8.175	1.00	46.44
20	ATOM	932	CE	LYS	A	95	169.767	37.553	8.161	1.00	48.82
20	ATOM	933	NZ	LYS	A	95	169.529	38.117	6.799	1.00	49.76
	ATOM	937	С	LYS	A	95	170.381	35.705	12.631	1.00	39.51
25	ATOM	938	0	LYS	A	95	170.781	36.869	12.594	1.00	39.68
	ATOM	939	N	ARG	A	96	171.000	34.743	13.304	1.00	39.70
30	ATOM	941	CA	ARG	A	96	172.229	34.965	14.065	1.00	38.81
30	ATOM	942	CB	ARG	A	96	173.386	35.432	13.159	1.00	38.58
	ATOM	943	CG	ARG	A	96	174.757	35.415	13.848	1.00	36.99
35	ATOM	944	CD	ARG	A	96	175.919	35.584	12.869	1.00	35.28
	MOTA	945	NE	ARG	A	96	177.207	35.640	13.562	1.00	37.59
40	MOTA	947	CZ	ARG	A	96	177.887	34.582	14.004	1.00	38.32
40	ATOM	948	NH1	ARG	A	96	177.425	33.351	13.832	1.00	41.87
	ATOM	951	NH2	ARG	A	96	179.024	34.758	14.658	1.00	34.85
45	MOTA	954	С	ARG	A	96	171.909	35.972	15.183	1.00	38.25
	MOTA	955	0	ARG	A	96	171.091	35.672	16.048	1.00	38.45
50	MOTA	956	N	PHE	A	97	172.506	37.161	15.158	1.00	36.05
	MOTA	958	CA	PHE	A	97	172.236	38.157	16.189	1.00	32.76
	MOTA	959	СВ	PHE	A	97	173.521	38.856	16.634	1.00	29.39
55	ATOM	960	CG	PHE	A	97	174.503	37.958	17.311	1.00	27.30
	ATOM	961	CD1	PHE	A	97	174.214	37.392	18.538	1.00	26.91
60	ATOM	962	CD2	ЬĤЕ	A	97	175.733	37.695	16.728	1.00	29.62
	ATOM	963	CE1	PHE	A	97	175.140	36.578	19.175	1.00	28.35
	MOTA	964	CE2	PHE	A	97	176.661	36.886	17.358	1.00	27.60
65	MOTA	965	CZ	PHE	A	97	176.364	36.327	18.582	1.00	28.30
	MOTA	966	С	PHE	A	97	171.261	39.208	15.693	1.00	34.25

	ATOM	967	0	PHE	A	97	171.016	40.199	16.387	1.00	33.16
5	MOTA	968	N	LEU	A	98	170.739	39.012	14.484	1.00	36.60
J	ATOM	970	CA	LEU	A	98	169.790	39.952	13.888	100	39.31
	MOTA	971	CB	LEU	A	98	169.874	39.905	12.361	1.00	38.74
10	ATOM	972	CG	LEU	A	98	169.875	41.238	11.608	1.00	38.17
	ATOM	973	CD1	LEU	A	98	169.883	40.951	10.128	1.00	40.67
15	MOTA	974	CD2	LEU	A	98	168.680	42.093	11.971	1.00	38.92
13	ATOM	975	С	LEU	A	98	168.366	39.631	14.317	1.00	40.51
	ATOM	976	0	LEU	A	98	167.854	38.554	14.016	1.00	42.93
20	ATOM	977	N	LYS	A	99	167.737	40.570	15.018	1.00	41.98
	ATOM	979	CA	LYS	A	99	166.365	40.410	15.488	1.00	42.37
25	ATOM	980	CB	LYS	A	99	166.258	40.705	16.989	1.00	41.90
23	ATOM	981	CG	LYS	A	99	166.690	39.566	17.898	1.00	45.06
	ATOM	982	CD	LYS	A	99	166.389	39.882	19.361	1.00	47.68
30	ATOM	983	CE	LYS	A	99	166.638	38.671	20.250	1.00	51.09
	ATOM	984	NZ	LYS	A	99	166.251	38.895	21.674	1.00	53.87
35	ATOM	988	С	LYS	A	99	165.440	41.351	14.727	1.00	43.86
	ATOM	989	0	LYS	A	99	165.634	42.573	14.739	1.00	45.41
	ATOM	990	N	MET	A	100	164.438	40.785	14.064	1.00	43.74
40	ATOM	992	CA	MET	A	100	163.479	41.575	13.310	1.00	43.78
	ATOM	993	CB	MET	A	100	163.576	41.234	11.826	1.00	44.48
45	ATOM	994	CG	MET	A	100	164.885	41.651	11.195	1.00	45.32
	MOTA	995	SD	MET	A	100	165.190	40.802	9.654	1.00	49.68
	ATOM	996	CE	MET	A	100	165.736	39.168	10.290	1.00	44.16
50	MOTA	997	С	MET	A	100	162.078	41.292	13.832	1.00	44.78
	MOTA	998	0	MET	A	100	161.693	40.132	13.987	1.00	44.97
55	ATOM	999	N	SER	A	101	161.335	42.352	14.140	1.00	46.57
	ATOM	1001	CA	SER	A	101	159.971	42.221	14.650	1.00	47.86
	MOTA	1002	CB	SER	Α	101	159.934	42.469	16.162	1.00	49.20
60	ATOM ·	1003	OG	SER	. A	101	160.560	43.693	16.509	1.00	51.70
	ATOM	1005	С	SER	A	101	159.009	43.168	13.937	1.00	48.68
65	MOTA	1006	0	SER	. A	101	159.428	44.161	13.332	1.00	48.84
	ATOM	1007	N	GLY	Α	102	157.722	42.839	13.986	1.00	49.46

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1.00 49.50 **GLY A 102** 156.716 43.664 13.343 MOTA 1009 CA 1.00 48.98 **GLY A 102** 155.359 42.989 13.341 C **ATOM** 1010 1.00 50.09 5 155.251 41.795 13.613 MOTA 1011 0 **GLY A 102** 1.00 48.68 MOTA 1012 N **ASN A 103** 154.317 43.766 13.079 1.00 47.35 ĆA **ASN A 103** 152.957 43.243 13.042 MOTA 1014 10 1.00 49.87 1015 CB **ASN A 103** 151.945 44.349 13.391 MOTA 152.061 44.837 14.840 1.00 51.69 CG **ASN A 103 ATOM** 1016 OD1 ASN A 103 45.423 1.00 53.03 153.067 15.234 15 1017 MOTA 1.00 53.17 ND2 ASN A 103 151.014 44.621 15.625 **ATOM** 1018 1.00 46.11 MOTA 1021 C **ASN A 103** 152.703 42.743 11.625 20 1.00 45.99 0 **ASN A 103** 153.459 43.073 10.708 MOTA 1022 PHE A 104 151.667 41.926 11.451 1.00 44.40 MOTA 1023 N 25 1.00 42.85 CA PHE A 104 151.306 41.410 10.131 ATOM 1025 PHE A 104 151.987 40.051 9.835 1.00 42.59 ATOM 1026 CB 38.849 10.482 1.00 41.53 1027 CG PHE A 104 151.319 MOTA 30 1.00 39.25 CD1 PHE A 104 151.650 38.458 11.776 ATOM 1028 1.00 39.79 CD2 PHE A 104 150.380 38.095 9.778 ATOM 1029 1.00 37.60 12.357 35 CE1 PHE A 104 151.058 37.339 ATOM 1030 1.00 36.72 36.976 CE2 PHE A 104 149.784 10.353 MOTA 1031 1.00 37.02 CZPHE A 104 150.125 36.599 11.644 MOTA 1032 40 149.788 1.00 42.22 ATOM 1033 C PHE A 104 41.303 10.006 PHE A 104 149.081 41.251 11.014 1.00 43.09 MOTA 1034 0 45 1.00 41.99 ATOM 1035 N **ASP A 105** 149.298 41.346 8.771 147.874 8.475 1.00 41.67 MOTA 1037 CA **ASP A 105** 41.227 MOTA 1038 CB **ASP A 105** 147.352 42,455 7.724 1.00 43.33 50 146.986 1.00 46.56 ATOM 1039 CG **ASP A 105** 43.602 8.648 OD1 ASP A 105 146.898 43.395 9.876 1.00 49.37 MOTA 1040 55 OD2 ASP A 105 146.770 44.720 8.141 1.00 51.38 MOTA 1041 1.00 41.95 MOTA 1042 C **ASP A 105** 147.658 39.985 7.622 0 **ASP A 105** 148.470 39.655 6.758 1.00 42.85 ATOM 1043 60 **LEU A 106** 146.538 7.841 1.00 40.99 MOTA 1044 N 39.322 1.00 39.85 ATOM CA LEU A 106 146.220 38.115 7.115 1046 65 1.00 38.87 MOTA 1047 CB **LEU A 106** 146.424 36.934 8.058 MOTA 1048 CG **LEU A 106** 146.224 35.489 7.629 1.00 39.36

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	ATOM	1049	CD1	LEU	A	106	146.825	34.605	8.690	1.00	40.16
5	ATOM	1050	CD2	LEU	Α	106	144.758	35.175	7.464	1.00	41.07
J	ATOM	1051	С	LEU	A	106	144.767	38.248	6.698	1.00	40.07
	MOTA	1052	0	LEU	A	106	143.968	38.841	7.414	1.00	41.85
10	ATOM	1053	N	SER	A	107	14.427	37.748	5.523	1.00	41.19
	ATOM	1055	CA	SER	A	107	143.054	37.830	5.061	1.00	43.48
15	ATOM	1056	CB	SER	A	107	142.911	38.907	3.979	1.00	45.09
13	ATOM	1057	OG	SER	A	107	143.440	40.155	4.402	1.00	50.35
	ATOM	1059	С	SER	A	107	142.633	36.484	4.502	1.00	43.83
20	MOTA	1060	0	SER	A	107	143.290	35.957	3.610	1.00	44.06
	ATOM	1061	N	ILE	A	108	141.587	35.900	5.077	1.00	45.70
25	MOTA	1063	CA	ILE	A	108	141.056	34.617	4.615	1.00	47.58
23	ATOM	1064	CB	ILE	A	108	140.829	33.644	5.789	1.00	46.97
	ATOM	1065	CG2	ILE	A	108	140.174	32.368	5.296	1.00	46.17
30	ATOM	1066	CG1	ILE	A	108	142.157	33.327	6.474	1.00	47.60
	ATOM	1067	CD1	ILE	A	108	142.042	32.368	7.637	1.00	49.72
35	MOTA	1068	С	ILE	A	108	139.716	34.922	3.959	1.00	48.97
55	MOTA	1069	0	ILE	A	108	138.818	35.433	4.619	1.00	51.76
	ATOM	.1070	N	GLU	A	109	139.582	34.652	2.664	1.00	50.18
40	MOTA	1072	CA	GLÜ	A	109	138.328	34.938	1.968	1.00	52.80
	MOTA	1073	CB	GLU	A	109	138.506	36.067	0.956	1.00	58.29
45	MOTA	1074	CG	GLU	A	109	138.871	37.409	1.556	1.00	67.61
	MOTA	1075	CD	GLU	Α	109	138.896	38.510	0.518	1.00	72.52
	MOTA	1076	OE1	GLU	A	109	139.591	38.345	-0.510	1.00	76.73
50	MOTA	1077	OE2	GLU	A	109	138.212	39.536	0.725	1.00	74.66
	ATOM	1078	С	GLU	A	109	137.771	33.733	1.244	1.00	51.51
55	MOTA	1079	0	GLU	A	109	138.449	32.717	1.091	1.00	51.13
•	MOTA	1080	N	GLY	A	110	136.539	33.867	0.767	1.00	50.94
	MOTA	1082	CA	GLY	A	110	135.904	32.778	0.051	1.00	49.27
60	ATOM	1083	С	GLY	A	110	135.714	31.551	0.920	1.00	48.08
	ATOM	1084	0	GLY	A	110	135.972	30.419	0.488	1.00	47.03
65	ATOM	1085	N	MET	A	111	135.318	31.784	2.168	1.00	46.98
	MOTA	1087	CA	MET	A	111	135.078	30.699	3.104	1.00	44.78

	MOTA	1088	CB	MET	A	111	135.376	31.143	4.536	1.00	44.81
	ATOM	1089	CG	MET	A	111	135.062	30.092	5.570	1.00	44.07
5	MOTA	1090	SD	MET	A	111	135.669	30.518	7.178	1.00	49.80
	ATOM	1091	CE	MET	A	111	136.833	29.192	7.434	1.00	46.06
10	ATOM	1092	C	MET	A	111	133.630	30.242	2.970	1.00	42.64
10	MOTA	1093	0	MET	A	111	132.712	31.061	2.960	1.00	40.20
٠. ر	MOTA	1094	N	SER	A	112	133.446	28.935	2.846	1.00	41.52
15	ATOM	1096	CA	SER	A	112	132.133	28.334	2.708	1.00	41.82
	MOTA	1097	СВ	SER	A	112	132.045	27.607	1.360	1.00	43.85
20	ATOM	1098	OG	SER	A	112	130.997	26.648	1.321	1.00	49.86
20	ATOM	1100	С	SER	A	112	131.908	27.363	3.861	1.00	41.93
	ATOM	1101	0	SER	A	112	132.725	26.467	4.097	1.00	41.84
25	ATOM	1102	N	ILE	A	113	130.816	27.569	4.592	1.00	42.07
	ATOM	1104	CA	ILE	A	113	130.444	26.722	5.725	1.00	40.13
30	ATOM	1105	CB	ILE	A	113	130.310	27.546	7.027	1.00	39.38
50	MOTA	1106	CG2	ILE	A	113	130.227	26.620	8.231	1.00	39.63
	ATOM	1107	CG1	ILE	A	113	131.505	28.480	7.190	1.00	35.78
35	ATOM	1108	CD1	ILE	A	113	131.284	29.547	8.225	1.00	38.78
	ATOM	1109	С	ILE	A	113	129.083	26.105	5.405	1.00	40.48
40	ATOM	1110	0	ILE	A	113	128.118	26.821	5.117	1.00	40.10
.0	MOTA	1111	N	SER	A	114	129.016	24.780	5.414	1.00	40.69
	MOTA	1113	CA	SER	A	114	127.775	24.076	5.126	1.00	39.86
45	ATOM	1114	CB	SER	A	114	127.915	23.277	3.835	1.00	39.27
	MOTA	1115	og	SER	A	114	126.695	22.645	3.486	1.00	45.23
50	MOTA	1117	С	SER	A	114	127.489	23.152	6.297	1.00	40.43
	MOTA	1118	0	SER	A	114	128.370	22.407	6.730	1.00	42.01
	MOTA	1119	N	ALA	A	115	126.261	23.198	6.804	1.00	40.94
55	MOTA	1121	CA	ALA	A	115	125.863	22.380	7.947	1.00	40.28
	MOTA	1122	CB	ALA	A	115	125.990	23.186	9.227	1.00	41.97
60	MOTA	1123	С	ALA	A	115	124.446	21.856	7.814	1.00	39.55
	ATOM	1124	0	ALA	A	115	123.551	22.581	7.385	1.00	38.65
	MOTA	1125	N .	ASP	A	116	124.248	20.603	8.213	1.00	39.92
65	ATOM	1127	CA	ASP	A	116	122.943	19.954	8.155	1.00	40.51
	MOTA	1128	CB	ASP	A	116	123.092	18.513	7.672	1.00	44.07

	ATOM	1129	CG	ASP A	116	123.359	18.421	6.184	1.00	48.91
5	ATOM	1130	OD1	ASP A	116	124.455	18.829	5.741	1.00	53.18
3	ATOM	1131	OD2	ASP A	116	122.467	17.940	5.455	1.00	52.27
	ATOM	1132	С	ASP A	116	122.241	19.956	9.504	1.00	39.36
10	ATOM	1133	0	ASP A	116	122.804	19.501	10.495	1.00	40.86
	ATOM	1134	N	LEU A	117	121.009	20.459	9.537	1.00	38.66
15	ATOM	1136	CA	LEU A	117	120.220	20.508	10.766	1.00	35.83
15	ATOM	1137	СВ	LEU A	. 117	119.578	21.888	10.944	1.00	34.68
	ATOM	1138	CG	LEU A	. 117	120.493	23.096	11.140	1.00	34.59
20	ATOM	1139	CD1	LEU A	. 117	119.667	24.361	11.279	1.00	35.67
	ATOM	1140	CD2	LEU A	. 117	121.346	22.895	12.370	1.00	35.88
25	MOTA	1141	С	LEU A	. 117	119.131	19.437	10.721	1.00	35.13
23	ATOM	1142	0	LEU A	117	118.287	19.435	9.824	1.00	35.15
	MOTA	1143	N	LYS A	118	119.155	18.531	11.691	1.00	35.41
30	ATOM	1145	CA	LYS A	118	118.180	17.450	11.776	1.00	34.03
	MOTA	1146	CB	LYS A	118	118.896	16.128	12.047	1.00	36.61
35	MOTA	1147	CG	LYS A	118	118.027	14.904	11.907	1.00	41.84
<i>J</i> J	MOTA	1148	CD	LYS A	118	118.870	13.640	11.890	1.00	45.83
	MOTA	1149	CE	LYS A	118	,117.998	12.400	11.773	1.00	49.70
40	MOTA	1150	NZ	LYS A	118	117.098	12.434	10.576	1.00	52.85
	ATOM	1154	С	LYS A	118	117.184	17.760	12.884	1.00	32.08
45	ATOM	1155	0	LYS A	118	117.572	18.045	14.014	1.00	30.72
	ATOM	1156	N	LEU A	119	115.900	17.737	12.545	1.00	34.05
	ATOM	1158	CA	LEU A	119	114.838	18.036	13.503	1.00	33.03
50	ATOM	1159	CB	LEU A	119	113.782	18.917	12.844	1.00	30.64
	ATOM	1160	CG	LEU A	119	114.277	20.278	12.372	1.00	26.32
55	ATOM	1161	CD1	LEU A	A 119	113.434	20.751	11.229	1.00	27.43
	ATOM	1162	CD2	LEU A	A 119	114.230	21.258	13.511	1.00	29.44
	MOTA	1163	С	LEU A	A 119	114.192	16.771	14.065	1.00	33.62
60	MOTA	1164	0	LEU A	A 119	113.904	15.819	13.334	1.00	32.69
	MOTA	1165	N	GLY I	À 120	113.952	16.776	15.368	1.00	33.78
65	MOTA	1167	CA	GLY 2	A 120	113.353	15.624	16.000	1.00	33.21
	ATOM	1168	С	GLY 3	A 120	112.431	16.059	17.109	1.00	34.60

	ATOM	1169	0	GLY	A	120	112.238	17.250	17.340	1.00	34.75
	MOTA	1170	N	SER	A	121	111.883	15.086	17.817	1.00	37.06
5	MOTA	1172	CA	SER	A	121	110.961	15.355	18.902	1.00	40.83
	MOTA	1173	СВ	SER	A	121	109.541	14.982	18.464	1.00	41.68
10	MOTA	1174	OG	SER	A	121	108.638	14.924	19.554	1.00	44.95
10	MOTA	1176	С	SER	A	121	111.357	14.527	20.108	1.00	43.32
	MOTA	1177	0	SER	A	121	111.990	13.476	19.975	1.00	44.77
15	MOTA	1178	N	ASN	A	122	111.034	15.038	21.287	1.00	45.49
	ATOM	1180	CA	ASN	A	122	111.308	14.330	22.525	1.00	48.51
20	ATOM	1181	СВ	ASN	A	122	112.014	15.244	23.523	1.00	47.83
20	MOTA	1182	CG	ASN	A	122	112.693	14.474	24.633	1.00	49.31
	ATOM	1183	OD1	ASN	A	122	112.445	13.283	24.827	1.00	48.55
25	MOTA	1184	ND2	ASN	A	122	113.574	15.146	25.359	1.00	52.06
	ATOM	1187	С	asn	A	122	109.925	13.955	23.040	1.00	50.18
30	MOTA	1188	0	ASN	A	122	109.197	14.810	23.534	1.00	51.08
50	MOTA	1189	N	PRO	A	123	109.530	12.678	22.895	1.00	51.66
	ATOM	1190	CD	PRO	A	123	110.378	11.593	22.375	1.00	51.97
35	ATOM	1191	CA	PRO	A	123	108.229	12.152	23.323	1.00	51.81
•	ATOM	1192	CB	PRO	A	123	108.291	10.696	22.875	1.00	52.33
40	ATOM	1193	CG	PRO	A	123	109.744	10.377	22.996	1.00	52.53
	MOTA	1194	С	PRO	A	123	107.903	12.255	24.808	1.00	52.83
	ATOM	1195	0	PRO	A	123	106.777	12.593	25.171	1.00	54.24
45	ATOM	1196	N	THR	A	124	108.879	11.969	25.664	1.00	53.70
	MOTA	1198	CA	THR	A	124	108.658	12.016	27.109	1.00	55.10
50	MOTA	1199	CB	THR	A	124	109.782	11.285	27.873	1.00	55.63
	MOTA	1200	OG1	THR	A	124	111.038	11.556	27.244	1.00	57.26
	ATOM	1202	CG2	THR	A	124	109.536	9.782	27.879	1.00	57.40
55	ATOM	1203	С	THR	A	124	108.467	13.418	27.687	1.00	54.39
	MOTA	1204	0	THR	A	124	108.031	13.568	28.833	1.00	56.00
60	MOTA	1205	N			125	108.794	14.439	26.901	1.00	52.41
	ATOM	1207	CA	SER	A	125	108.647	15.824	27.345	1.00	49.57
	MOTA	1208	CB			125	110.023	16.448	27.598	1.00	49.34
65	ATOM	1209	OG	SER	A	125	110.859	16.328	26.459	1.00	50.88
	MOTA	1211	С	SER	A	125	107.855	16.676	26.355	1.00	46.68

	ATOM	1212	0	SER A 125	107.366	17.747	26.702	1.00 46.69
_	ATOM	1213	N	GLY A 126	107.734	16.191	25.123	1.00 44.84
5	ATOM	1215	CA	GLY A 126	107.008	16.906	24.091	1.00 42.00
	ATOM	1216	С	GLY A 126	107.739	18.113	23.537	1.00 40.90
10	ATOM	1217	0	GLY A 126	107.125	18.953	22.885	1.00 41.68
	ATOM	1218	N	LYS A 127	109.044	18.201	23.780	1.00 40.27
1.5	ATOM	1220	CA	LYS A 127	109.846	19.330	23.303	1.00 38.06
15	ATOM	1221	СВ	LYS A 127	110.914	19.708	24.340	1.00 41.48
	ATOM	1222	CG	LYS A 127	110.408	20.194	25.705	1.00 44.59
20	MOTA	1223	CD	LYS A 127	109.966	21.657	25.697	1.00 48.40
	ATOM	1224	CE	LYS A 127	109.668	22.142	27.118	1.00 49.68
25	MOTA	1225	NZ	LYS A 127	109.045	23.498	27.151	1.00 50.76
25	MOTA	1229	С	LYS A 127	110.533	18.984	21.980	1.00 36.38
	MOTA	1230	0	LYS A 127	110.916	17.827	21.751	1.00 36.93
30	ATOM	1231	N	PRO A 128	110.705	19.980	21.093	1.00 33.49
	MOTA	1232	CD	PRO A 128	110.310	21.395	21.226	1.00 31.99
35	ATOM	1233	CA	PRO A 128	111.354	19.747	19.803	1.00 31.41
33	ATOM	1234	CB	PRO A 128	111.023	21.020	19.037	1.00 31.27
	ATOM	1235	CG	PRO A 128	111.085	22.056	20.109	1.00 29.23
40	MOTA	1236	С	PRO A 128	112.853	19.611	20.007	1.00 31.58
	ATOM	1237	0	PRO A 128	113.389	20.080	21.010	1.00 31.45
45	MOTA	1238	N	THR A 129	113.523	18.936	19.087	1.00 31.21
	MOTA	1240	CA	THR A 129	114.960	18.779	19.181	1.00 30.79
	ATOM	1241	CB	THR A 129	115.377	17.332	19.510	1.00 29.38
50	MOTA	1242	OG1	THR A 129	114.952	16.451	18.467	1.00 30.58
	MOTA	1244	CG2	2 THR A 129	114.773	16.886	20.818	1.00 27.50
55	MOTA	1245	С	THR A 129	115.568	19.185	17.856	1.00 32.76
	ATOM	1246	0	THR A 129	114.907	19.148	16.813	1.00 32.88
	MOTA	1247	N	ILE A 130	116.828	19.582	17.904	1.00 35.82
60	ATOM	1249	CA	ILE A 130	117.554	19.999	16.717	1.00 37.64
	MOTA	1250	CB.	ILE A 130	117.302	21.506	16.401	1.00 38.78
65	ATOM	1251	CG2	2 ILE A 130	117.277	22.335	17.665	1.00 40.02
95	ATOM	1252	CG:	L ILE A 130	118.345	22.042	15.425	1.00 38.94

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	MOTA	1253	CD1	ILE	A	130	118.138	21.591	14.014	1.00	42.34
	ATOM	1254	С	ILE	A	130	119.026	19.740	16.985	1.00	38.69
5	MOTA	1255	0	ILE	A	130	119.534	20.085	18.048	1.00	40.22
	MOTA	1256	N	THR	A	131	119.681	19.039	16.069	1.00	39.62
10	MOTA	1258	CA	THR	A	131	121.098	18.748	16.217	1.00	40.39
10	MOTA	1259	СВ	THR	A	131	121.351	17.318	16.747	1.00	40.39
	MOTA	1260	OG1	THR	A	131	120.833	16.354	15.825	1.00	40.07
15	MOTA	1262	CG2	THR	A	131	120.696	17.124	18.113	1.00	43.21
	ATOM	1263	С	THR	A	131	121.788	18.927	14.878	1.00	41.17
20	ATOM	1264	0	THR	A	131	121.139	18.951	13.839	1.00	41.86
20	MOTA	1265	N	CYS	À	132	123.099	19.107	14.911	1.00	42.72
	MOTA	1267	CA	CYS	A	132	123.875	19.284	13.696	1.00	42.81
25	ATOM '	1268	CB	CYS	A	132	124.963	20.326	13.932	1.00	42.56
	ATOM	1269	SG	CYS	A	132	126.009	20.629	12.530	1.00	38.05
30	MOTA	1270	С	CYS	A	132	124.494	17.940	13.333	1.00	44.48
	MOTA	1271	0	CYS	A	132.	125.370	17.446	14.041	1.00	46.08
	MOTA	1272	N	SER	A	133	124.019	17.344	12.244	1.00	45.05
35	ATOM	1274	CA	SER	A	133	124.508	16.045	11.794	1.00	45.93
	MOTA	1275	CB	SER	A	133	123.410	15.315	11.015	1.00	47.62
40	MOTA	1276	OG	SER	A	133	122.962	16.082	9.908	1.00	51.90
	ATOM	1278	С	SER	A	133 '	125.793	16.072	10.966	1.00	45.39
	MOTA	1279	0	SER	A	133	126.512	15.078	10.916	1.00	46.00
45	MOTA	1280	N	SER	A	134	126.056	17.182	10.282	1.00	45.72
	MOTA	1282	CA	SER	A	134	127.257	17.322	9.453	1.00	45.69
50	MOTA	1283	CB	SER	A	134	127.048	16.672	8.073	1.00	47.17
	MOTA	1284	OG	SER	A	134	127.221	15.263	8.110	1.00	50.20
	ATOM	1286	С	SER	A	134	127.642	18.788	9.265	1.00	44.02
55	MOTA	1287	0	SER	A	134	126.775	19.663	9.212	1.00	45.04
	MOTA	1288	N	CYS	A	135	128.939	19.047	9.145	1.00	42.35
60	MOTA	1290	CA	CYS	A	135	129.442	20.398	8.941	1.00	40.67
	ATOM	1291	С	CYS	A	135	130.728	20.345	8.144	1.00	41.67
	ATOM	1292	0	CYS	A	135	131.469	19.361	8.216	1.00	40.71
65	ATOM	1293	СВ	CYS	A	135	129.718	21.084	10.274	1.00	39.61
	MOTA	1294	SG	CYS	A	135	130.296	22.800	10.094	1.00	34.66

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	MOTA	1295	N	SER A	136	130.975	21.383	7.354	1.00	43.65
5	MOTA	1297	CA	SER A	136	132.192	21.451	6.562	1.00	45.71
5	MOTA	1298	СВ	SER A	136	132.131	20.504	5.357	1.00	48.55
	ATOM	1299	OG	SER A	136	131.231	20.961	4.363	1.00	54.17
10	MOTA	1301	С	SER A	A 136	132.465	22.874	6.114	1.00	45.23
	ATOM	1302	0	SER A	136	131.582	23.551	5.583	1.00	44.66
15	MOTA	1303	N	SER A	137	133.673	23.339	6.411	1.00	45.29
15	ATOM	1305	CA	SER A	137	134.110	24.671	6.040	1.00	45.90
	MOTA	1306	CB	SER A	137	134.896	25.302	7.186	1.00	45.32
20	ATOM	1307	OG	SER A	137	135.302	26.620	6.867	1.00	46.54
	ATOM	1309	С	SER A	137	134.990	24.514	4.806	1.00	47.49
25	MOTA	1310	0	SER A	137	135.330	23.392	4.416	1.00	48.14
23	MOTA	1311	N	HIS A	138	135.371	25.630	4.200	1.00	48.87
	ATOM	1313	CA	HIS A	138	136.196	25.599	3.003	1.00	49.64
30	MOTA	1314	CB	HIS A	138	135.338	25.149	1.811	1.00	53.41
	MOTA	1315	CG	HIS A	138	136.097	24.992	0.529	1.00	56.80
35	MOTA ·	1316	CD2	HIS A	A 138	136.712	23.915	-0.016	1.00	58.14
33	MOTA	1317	ND1	HIS A	A 138	136.268	26.025	-0.368	1.00	59.15
	MOTA	1319	CE1	HIS A	A 138	136.956	25.592	-1.412	1.00	59.11
40	MOTA	1320	NE2	HIS A	A 138	137.238	24.316	-1.222	1.00	59.30
	MOTA	1322	С	HIS A	A 138	136.779	26.983	2.742	1.00	49.79
45	MOTA	1323	0	HIS A	A 138	136.042	27.938	2.510	1.00	49.66
	MOTA	1324	N	ILE A	A 139	138.102	27.082	2.808	1.00	50.32
	MOTA	1326	CA	ILE A	A 139	138.822	28.330	2.565	1.00	50.43
50	MOTA	1327	CB	ILE A	A 139	139.937	28.519	3.606	1.00	50.04
	MOTA	1328	CG2	ILE A	A 139	140.876	29.651	3.195	1.00	46.81
55	MOTA	1329	CG1	ILE A	A 139	139.313	28.751	4.982	1.00	49.41
	MOTA	1330	CD1	ILE A	A 139	140.310	28.816	6.105	1.00	53.48
	MOTA	1331	С	ILE A	A 139	139.430	28.312	1.159	1.00	51.98
60	ATOM	1332	0	ILE A	A 139	140.019	27.311	0.743	1.00	51.75
	MOTA	1333	N .	ASN A	A 140	139.267	29.409	0.425	1.00	53.16
65	MOTA	1335	CA	ASN I	A 140	139.792	29.503	-0.935	1.00	54.65
	MOTA	1336	CB	ASN I	A 140	138.824	30.291	-1.832	1.00	57.65

	ATOM	1337	CG	ASN A	140	139.311	30.400	-3.276	1.00	60.64"
	ATOM	1338	OD1	ASN A I	L40	139.295	31.483	-3.867	1.00	61.18
5	ATOM	1339	ND2	ASN A 1	140	139.742	29.279	-3.848	1.00	62.27
	MOTA	1342	С	ASN A	140	141.203	30.092	-1.024	1.00	53.67
10	MOTA	1343	0	ASN A	140	142.097	29.463	-1.589	1.00	53.36
10	ATOM	1344	N	SER A	141	141.401	31.289	-0.474	1.00	52.97
	ATOM	1346	CA	SER A I	141	142.711	31.943	-0.518	1.00	52.86
15	MOTA	1347	СВ	SER A 1	141	142.728	33.051	-1.585	1.00	52.81
	ATOM	1348	OG	SER A	141	141.928	34.167	-1.218	1.00	50.44
20	MOTA	1350	C	SER A	141	143.114	32.526	0.832	1.00	52.49
20	MOTA	1351	0	SER A	141	142.263	32.778	1.684	1.00	54.02
	MOTA	1352	N .	VAL A	142	144.415	32.724	1.023	1.00	52.25
25	MOTA	1354	CA	VAL A	142	144.947	33.290	2.256	1.00	50.52
	MOTA	1355	СВ	VAL A	142	145.583	32.203	3.152	1.00	49.93
30	MOTA	1356	CG1	VAL A	142	146.214	32.826	4.380	1.00	49.68
30	MOTA	1357	CG2	VAL A	142	144.533	31.197	3.577	1.00	50.61
	MOTA	1358	С	VAL A	142	145.990	34.335	1.880	1.00	51.54
35	MOTA	1359	0	VAL A	142	147.083	34.000	1.418	1.00	50.89
	MOTA	1360	N	HIS A	143	145.611	35.603	2.021	1.00	54.44
40	MOTA	1362	CA	HIS A	143	146.470	36.742	1.709	1.00	57.63
	MOTA	1363	CB	HIS A	143	145.639	37.904	1.154	1.00	60.05
	MOTA	1364	CG	HIS A	143	145.323	37.789	-0.306	1.00	65.61
45	ATOM	1365	CD2	HIS A	143	145.600	38.615	-1.343	1.00	67.19
	ATOM	1366	ND1	HIS A	143	144.632	36.722	-0.841	1.00	67.65
50	ATOM	1368	CE1	HIS A	143	144.498	36.896	-2.145	1.00	68.08
	MOTA	1369	NE2	HIS A	143	145.078	38.037	-2.473	1.00	67.35
	MOTA	1371	С	HIS A	143	147.229	37.232	2.936	1.00	58.93
55	MOTA	1372	0	HIS A	143	146.663	37.914			58.93
	MOTA	1373	N	VAL A		148.510	36.893	3.012	1.00	60.40
60	MOTA	1375	CA	VAL A	144	149.347	37.316	4.124	1.00	62.35
	MOTA	1376	CB	VAL A	144	150.423	36.270	4.441	1.00	61.34
	ATOM	1377	CG1	VAL A	144	151.245	36.703	5.641	1.00	62.08
65	MOTA	1378	CG2	VAL A	144	149.771	34.938	4.709	1.00	62.01
	MOTA	1379	С	VAL A	144	150.000	38.632	3.724	1.00	64.53

	ATOM	1380	0	VAL A 144	150.971	38.658	2.971	1.00	64.35
5	MOTA	1381	N	HIS A 145	149.440	39.729	4.214	1.00	67.48
3	ATOM	1383	CA	HIS A 145	149.945	41.050	3.886	1.00	71.08
	ATOM	1384	СВ	HIS A 145	148.791	42.045	3.771	1.00	72.13
10	MOTA	1385	CG	HIS A 145	147.860	41.751	2.642	1.00	74.06
	ATOM	1386	CD2	HIS A 145	148.098	41.536	1.327	1.00	74.26
15	ATOM	1387	ND1	HIS A 145	146.497	41.640	2.810	1.00	75.07
13	ATOM	1389	CE1	HIS A 145	145.934	41.370	1.646	1.00	75.64
	ATOM	1390	NE2	HIS A 145	146.884	41.303	0.730	1.00	75.46
20	ATOM	1392	С	HIS A 145	150.998	41.600	4.830	1.00	73.50
	ATOM	1393	0	HIS A 145	150.742	42.560	5.553	1.00	74.03
25	ATOM	1394	N	ILE A 146	152.184	40.999	4.826	1.00	76.22
23	MOTA	1396	CA	ILE A 146	153.277	41.497	5.660	1.00	78.65
	ATOM	1397	CB	ILE A 146	154.282	40.374	6.055	1.00	77.20
30	ATOM	1398	CG2	ILE A 146	155.320	40.905	7.037	1.00	75.80
	ATOM	1399	CG1	ILE A 146	153.545	39.214	6.729	1.00	76.58
35	ATOM	1400	CD1	ILE A 146	154.451	38.091	7.184	1.00	75.69
	ATOM	1401	С	ILE A 146	153.972	42.574	4.815	1.00	81.85
	MOTA	1402	0	ILE A 146	155.107	42.970	5.082	1.00	81.33
40	ATOM	1403	N	SER A 147	153.273	43.023	3.774	1.00	85.84
	MOTA	1405	CA	SER A 147	153.758	44.049	2.864	1.00	89.73
45	ATOM	1406	СВ	SER A 147	152.736	44.267	1.737	1.00	89.11
	ATOM	1407	OG	SER A 147	152.396	43.046	1.097	1.00	89.21
	ATOM	1409	С	SER A 147	153.983	45.354	3.637	1.00	92.25
50	ATOM	1410	0	SER A 147	153.080	46.188	3.753	1.00	94.01
	ATOM	1411	N	ALA A 148	155.182	45.494	4:197	1.00	92.99
55	ATOM	1413	CA	ALA A 148	155.568	46.675	4.967	1.00	92.04
	ATOM	1414	CB	ALA A 148	155.027	46.577	6.395	1.00	92.57
	ATOM	1415	С	ALA A 148	157.092	46.778	4.982	1.00	91.03
60	ATOM	1416	0	ALA A 148	157 .657	47.869	5.086	1.00	92.24
	ATOM	1417	N	ALA A 149	157.749	45.628	4.875	1.00	88.25
65	MOTA	1419	CA	ALA A 149	159.201	45.558	4.864	1.00	85.00
	MOTA	1420	CB	ALA A 149	159.720	45.237	6.263	1.00	85.37

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	ATOM	1421	С	ALA A 149	159.624	44.474	3.874	1.00 82.20-
	ATOM	1422	0	ALA A 149	158.888	43.508	3.650	1.00 83.34
5	ATOM	1423	N	SER A 150	160.785	44.657	3.252	1.00 77.07
	MOTA	1425	CA	SER A 150	161.304	43.690	2.293	1.00 71.37
10	ATOM	1426	СВ	SER A 150	162.463	44.313	1.509	1.00 72.17
10	ATOM	1427	OG	SER A 150	162.092	45.574	0.968	1.00 72.35
	MOTA	1429	С	SER A 150	161.768	42.434	3.037	1.00 66.95
15	MOTA	1430	0	SER A 150	162.927	42.330	3.440	1.00 66.96
٠	ATOM	1431	N	VAL A 151	160.847	41.496	3.240	1.00 60.88
20	MOTA	1433	CA	VAL A 151	161.146	40.257	3.949	1.00 54.82
20	ATOM	1434	СВ	VAL A 151	160.736	40.381	5.446	1.00 55.79
	MOTA	1435	CG1	VAL A 151	159.233	40.593	5.584	1.00 55.13
25	MOTA	1436	CG2	VAL A 151	161.206	39.176	6.242	1.00 55.16
	MOTA	1437	С	VAL A 151	160.417	39.104	3.259	1.00 51.49
30	ATOM	1438	0	VAL A 151	159.721	38.308	3.887	1.00 51.42
50	MOTA	1439	N	GLY A 152	160.639	38.992	1.957	1.00 48.87
	ATOM	1441	CA	GLY A 152	159.994	37.961	1.167	1.00 45.39
35	MOTA	1442	С	GLY A 152	160.100	36.531	1.656	1.00 42.87
	MOTA	1443	0	GLY A 152	159.144	35.766	1.543	1.00 43.01
40	MOTA	1444	N	TRP A 153	161.245	36.160	2.211	1.00 41.58
-	MOTA	1446	CA	TRP A 153	161.432	34.795	2.682	1.00 41.92
	MOTA	1447	CB	TRP A 153	162.853	34.591	3.196	1.00 42.78
45	MOTA	1448	CG	TRP A 153	163.153	35.267	4.501	1.00 43.10
	MOTA	1449	CD2	TRP A 153	163.738	36.559	4.678	1.00 43.00
50	MOTA	1450	CE2	TRP A 153	163.953	36.738	6.064	1.00 43.32
	MOTA	1451	CE3	TRP A 153	164.112	37.583	3.796	1.00 45.47
	MOTA	1452	CD1	TRP A 153	163.022	34.730	5.760	1.00 42.27
- 55	ATOM	1453	NEl	TRP A 153	163.508	35.607	6.700	1.00 42.43
	MOTA	1455	CZ2	TRP A 153	164.528	37.897	6.587	1.00 44.70
60	MOTA	1456	CZ3	TRP A 153	164.686	38.738	4.317	1.00 47.45
	MOTA	1457	CH2	TRP A 153	164.889	38.885	5.701	1.00 47.30
	ATOM	1458	С	TRP A 153	160.442	34.393	3.763	1.00 41.11
65	ATOM	1459	0	TRP A 153	.160.053		3.852	1.00 41.06
	MOTA	1460	И	LEU A 154	160.045	35.357	4.585	1.00 40.39

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	MOTA	1462	CA	LEU A	154	159.109	35.103	5.669	1.00 38.88
5	MOTA	1463	CB	LEU A	154	159.080	36.284	6.618	1.00 37.14
3	MOTA	1464	CG	LEU A	154	158.607	35.931	8.011	1.00 35.27
	MOTA	1465	CD1	LEU A	154	159.491	34.860	8.602	1.00 35.45
10	ATOM	1466	CD2	LEU A	154	158.655	37.166	8.838	1.00 37.10
	ATOM	1467	C	LEU A	154	157.721	34.847	5.115	1.00 38.87
15	ATOM	1468	0	LEU A	154	156.997	33.985	5.611	1.00 39.93
13	ATOM	1469	N	ILE A	155	157.357	35.596	4.079	1.00 40.01
	ATOM	1471	CA	ILE A	155	156.062	35.435	3.430	1.00 40.45
20	ATOM	1472	CB	ILE A	155	155.765	36.579	2.454	1.00 39.45
	ATOM	1473	CG2	ILE A	155	154.467	36.307	1.721	1.00 39.70
25	ATOM	1474	CG1	ILE A	155	155.679	37.906	3.215	1.00 39.53
23	MOTA	1475	CD1	ILE A	155	155.426	39.114	2.337	1.00 39.05
	ATOM	1476	С	ILE A	155	156.048	34.106	2.686	1.00 42.61
30	MOTA	1477	0	ILE A	155	155.015	33.435	2.619	1.00 44.53
	MOTA	1478	N	GLN A	156	157.187	33.745	2.100	1.00 43.49
35	MOTA	1480	CA	GLN A	156	157.323	32.475	1.398	1.00 44.17
	MOTA	1481	CB	GLN A	156	158.708	32.368	0.760	1.00 48.67
	MOTA	1482	CG	GLN A	156	159.097	30.954	0.310	1.00 55.67
40	MOTA	1483	CD	GLN A	156	160.369	30.440	0.988	1.00 59.91
	MOTA	1484	OE1	GLN A	156	160.673	30.802	2.125	1.00 61.96
45	MOTA	1485	NE2	GLN A	156	161.114	29.593	0.285	1.00 60.94
	MOTA	1488	С	GLN A	156	157.163	31.384	2.452	1.00 43.64
	MOTA	1489	0	GLN A	156	156.445	30.408	2.249	1.00 43.84
50	MOTA	1490	И	LEU A	157	157.834	31.581	3.585	1.00 43.44
	MOTA	1492	CA	LEU A	. 157	157.794	30.653	4.704	1.00 41.27
55	MOTA	1493	CB		157	158.644	31.175	5.866	1.00 42.44
	MOTA	1494	CG	LEU A	. 157	160.152	30.930	5.797	1.00 42.39
	MOTA	1495	CD1	LEU A	157	160.873	31.722	6.876	1.00 41.73
60	MOTA	1496	CD2	LEU A		160.426	29.447	5.948	1.00 42.15
	ATOM	1497	С		157	156.372	30.423	5.173	1.00 40.27
65	· MOTA	1498	0	LEU A	157	155.948	29.280	5.328	1.00 41.05
	MOTA	1499	N	PHE A	158	155.620	31.497	5.378	1.00 39.18

	ATOM	1501	CA	PHE 2	A	158	154.246	31.341	5.821	1.00	41.38
	MOTA	1502	СВ	PHE I	A	158	153.519	32.684	5.927	1.00	40.94
5	MOTA	1503	CG	PHE 2	A	158	152.145	32.570	6.537	1.00	43.98
	MOTA	1504	CD1	PHE 2	A	158	151.068	32.087	5.788	1.00	44.07
10	ATOM	1505	CD2	PHE 2	A	158	151.939	32.886	7.878	1.00	43.45
10	MOTA	1506	CE1	PHE :	A	158	149.814	31.915	6.367	1.00	42.96
	MOTA	1507	CE2	PHE .	A	158	150.690	32.718	8.465	1.00	41.63
15	MOTA	1508	CZ	PHE .	A	158	149.626	32.231	7.707	1.00	43.20
	MOTA	1509	С	PHE .	A	158	153.472	30.409	4.893	1.00	42.96
20	MOTA	1510	0	PHE	A	158	152.862	29.443	5.350	1.00	44.53
20	ATOM	1511	N	HIS .	A	159	153.513	30.682	3.594	1.00	45.12
	MOTA	1513	CA	HIS	A	159	152.790	29.860	2.624	1.00	46.02
25	ATOM	1514	СВ	HIS	A	159	152.808	30.507	1.230	1.00	45.21
	ATOM	1515	CG	HIS	A	159	152.049	31.796	1.145	1.00	44.03
30	MOTA	1516	CD2	HIS	A	159	152.477	33.081	1.099	1.00	44.13
30	ATOM	1517	ND1	HIS	A	159	150.674	31.851	1.087	1.00	44.17
	ATOM	1519	CE1	HIS	A	159	150.285	33.111	1.012	1.00	45.07
35	MOTA	1520	NE2	HIS	A	159	151.360	33.878	1.016	1.00	45.02
	MOTA	1522	С	HIS	A	159	153.314	28.426	2.524	1.00	47.97
40	MOTA	1523	0	HIS	A	159	152.530	27.491	2.353	1.00	48.86
70	MOTA	1524	N	LYS	A	160	154.624	28.253	2.678	1.00	49.77
	MOTA	1526	CA	LYS	A	160	155.246	26.935	2.562	1.00	50.79
45	MOTA	1527	CB	LYS	A	160	156.664	27.080	1.990	1.00	51.90
	MOTA	1528	CG	LYS	A	160	157.285	25.768	1.539	1.00	53.77
50	MOTA	1529	CD	LYS	A	160	158.669	25.957	0.950	1.00	54.99
	ATOM	1530	CE	LYS	A	160	159.130	24.690	0.231	1.00	57.16
	MOTA	1531	NZ	LYS	A	160	158.253	24.330	-0.934	1.00	55.19
55	MOTA	1535	С	LYS	A	160	155.286	26.065	3.825	1.00	51.12
	MOTA	1536	0	LYS	A	160.	155.537	24.862	3.736	1.00	52.93
60	ATOM	1537	N	LYS	A	161	155.031	26.646	4.991	1.00	49.90
	MOTA	1539	CA	LYS	A	161	155.086	25.873	6.229	1.00	48.78
	ATOM	1540	СВ	LYS	A	161	156.340	26.263	7.016	1.00	51.36
65	MOTA	1541	CG	LYS	A	161	157.660	25.983	6.310	1.00	55.24
	ATOM	1542	CD	LYS	A	161	158.186	24.592	6.626	1.00	59.40

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	MOTA	1543	CE	LYS A 161	158.484	24.443	8.114	1.00 61.04
_	ATOM.	1544	NZ	LYS A 161	159.015	23.095	8.464	1.00 62.60
5	MOTA	1548	С	LYS A 161	153.875	26.012	7.149	1.00 47.20
	MOTA	1549	0	LYS A 161	153.460	25.042	7.787	1.00 46.35
10	MOTA	1550	N	ILE A 162	153.311	27.214	7.209	1.00 46.07
	ATOM	1552	CA	ILE A 162	152.186	27.506	8.094	1.00 45.42
	ATOM	1553	CB	ILE A 162	152.421	28.854	8.806	1.00 45.05
15	MOTA	1554	CG2	ILE A 162	151.211	29.251	9.646	1.00 45.73
	MOTA	1555	CG1	ILE A 162	153.679	28.764	9.670	1.00 43.35
20	MOTA	1556	CD1	ILE A 162	154.015	30.042	10.353	1.00 43.37
	MOTA	1557	С	ILE A 162	150.764	27.483	7.520	1.00 45.58
25	MOTA	1558	0	ILE A 162	149.849	26.960	8.157	1.00 45.52
25	MOTA	1559	N	GLU A 163	150.570	28.036	6.329	1.00 46.35
	MOTA	1561	CA	GLU A 163	149.239	28.093	5.731	1.00 47.82
30	MOTA	1562	СВ	GLU A 163	149.310	28.624	4.302	1.00 49.25
	ATOM	1563	CG	GLU A 163	147.956	29.028	3.735	1.00 51.68
35	MOTA	1564	CD	GLU A 163	147.965	29.204	2.224	1.00 52.54
33	MOTA	1565	OE1	GLU A 163	149.007	29.597	1.658	1.00 53.57
	MOTA	1566	OE2	GLU A 163	146.918	28.941	1.599	1.00 53.52
40	MOTA	1567	C	GLU A 163	148.470	26.775	5.746	1.00 47.94
	MOTA	1568	0	GLU A 163	147.289	26.757	6.069	1.00 49.00
45	ATOM	1569	N	SER A 164	149.142	25.678	5.420	1.00 49.09
75	MOTA	1571	CA	SER A 164	148.510	24.360	5.383	1.00 49.75
	ATOM	1572	CB	SER A 164	149.486	23.327	4.818	1.00 50.94
50	ATOM	1573	OG	SER A 164	149.971	23.737	3.548	1.00 56.63
	MOTA	1575	C	SER A 164	147.953	23.865	6.717	1.00 49.89
55	ATOM	1576	0	SER A 164	147.100	22.982	6.739	1.00 52.82
20	MOTA	1577	N	ALA A 165	148.442	24.415	7.824	1.00 48.05
	MOTA	1579	CA	ALA A 165	147.978	24.017	9.151	1.00 44.58
60	MOTA	1580	СВ	ALA A 165	149.153	23.920	10.109	1.00 45.11
	MOTA	1581	С	ALA A 165	146.947	25.009	9.678	1.00 43.87
65	ATOM	1582	.0	ALA A 165	145.980	24.626	10.335	1.00 43.62
0.5	MOTA	1583	N	LEU A 166	147.169	26.287	9.391	1.00 43.88

	MOTA	1585	CA	LEU A	166	146.267	27.349	9.812	1.00	44.57
	MOTA	1586	СВ	LEU A	166	146.783	28.701	9.313	1.00	41.17
5	MOTA	1587	CG	LEU A	166	145.977	29.957	9.655	1.00	39.43
	MOTA	1588	CD1	LEU A	166	146.438	30.508	10.979	1.00	40.32
10	MOTA	1589	CD2	LEU A	166	146.165	31.005	8.577	1.00	39.33
10	MOTA	1590	С	LEU A	166	144.870	27.094	9.246	1.00	47.80
	ATOM	1591	0	LEU A	166	143.899	27.003	9.995	1.00	50.31
15	MOTA	1592	N	ARG A	167	144.777	26.946	7.927	1.00	49.83
	MOTA	1594	CA	ARG A	167	143.493	26.714	7.274	1.00	53.68
20	MOTA	1595	CB	ARG A	167	143.659	26.564	5.753	1.00	54.31
20	MOTA	1596	CG	ARG A	167	144.488	25.354	5.329	1.00	57.76
	MOTA	1597	CD	ARG A	167	144.282	24.966	3.869	1.00	58.03
25	MOTA	1598	NE	ARG A	167	144.657	26.032	2.948	1.00	56.39
	MOTA	1600	CZ	ARG A	167	143.786	26.753	2.256	1.00	55.99
30	ATOM	1601	NHl	ARG A	167	142.485	26.525	2.380	1.00	56.36
	MOTA	1604	NH2	ARG A	167	144.214	27.699	1.436	1.00	55.25
	ATOM	1607	С	ARG A	167	142.774	25.495	7.836	1.00	56.26
35	ATOM	1608	0	ARG A	167	141.570	25.544	8.088	1.00	58.47
	ATOM	1609	N	ASN A	168	143.517	24.418	8.077	1.00	59.06
40	ATOM	1611	CA	ASN A	168	142.924	23.188	8.596	1.00	60.42
.0	ATOM	1612	CB	ASN A	168	143.936	22.037	8.586	1.00	63.69
	ATOM	1613	CG	asn a	168	143.264	20.670	8.594	1.00	69.44
45	MOTA	1614	OD1	ASN A	168	142.186	20.491	8.021	1.00	73.41
	MOTA	1615	ND2	ASN A	168	143.902	19.696	9.232	1.00	71.02
50	ATOM	1618	С	ASN A	168	142.358	23.378	9.995	1.00	58.54
	MOTA	1619	0	ASN A	168	141.291	22.857	10.312	1.00	59.34
	MOTA	1620	N	LYS A	169	143.059	24.138	10.826	1.00	56.61
55	MOTA	1622	CA	LYS A	169	142.591	24.380	12.179	1.00	54.57
	MOTA	1623	СВ	LYS A	169	143.732	24.882	13.065	1.00	56.85
60	MOTA	1624	CG	LYS A	169	143.365	25.002	14.540	1.00	60.65
00	MOTA	1625	CD	LYS A	169	144.599	24.927	15.425	1.00	65.37
	MOTA	1626	CE	LYS A	169	145.276	23.560	15.329	1.00	68.82
65	MOTA	1627	NZ	LYS A	169	146.576	23.504	16.062	1.00	70.63
	MOTA	1631	С	LYS A	169	141.439	25.372	12.173	1.00	51.59

	MOTA	1632	0	LYS A 169	140.570	25.327	13.035	1.00 51.73
5	ATOM	1633	N	MET A 170	141.426	26.264	11.192	1.00 49.16
3	ATOM	1635	CA	MET A 170	140.362	27.251	11.090	1.00 47.96
	MOTA	1636	СВ	MET A 170	140.733	28.340	10.087	1.00 49.44
10	MOTA	1637	CG	MET A 170	139.792	29.528	10.105	1.00 52.89
	ATOM	1638	SD	MET A 170	139.730	30.289	11.734	1.00 55.23
1.5	ATOM	1639	CE	MET A 170	141.170	31.310	11.695	1.00 55.26
15	ATOM	1640	С	MET A 170	139.075	26.565	10.659	1.00 45.37
	ATOM	1641	0	MET A 170	137.989	26.892	11.139	1.00 46.51
20	ATOM	1642	N	ASN A 171	139.207	25.601	9.759	1.00 42.65
	ATOM	1644	CA	ASN A 171	138.063	24.858	9.268	1.00 41.99
25	ATOM	1645	СВ	ASN A 171	138.448	24.034	8.039	1.00 42.92
25	ATOM	1646	CG	ASN A 171	138.644	24.889	6.795	1.00 43.47
	ATOM	1647	OD1	ASN A 171	138.041	25.957	6.658	1.00 44.24
30	ATOM	1648	ND2	ASN A 171	139.478	24.415	5.875	1.00 44.77
	MOTA	1651	С	ASN A 171	137.487	23.962	10.353	1.00 42.70
35	MOTA	1652	0	ASN A 171	136.284	23.701	10.374	1.00 43.68
33	ATOM	1653	N	SER A 172	138.341	23.502	11.263	1.00 44.20
	MOTA	1655	CA	SER A 172	137.898	22.645	12.358	1.00 45.11
40	MOTA	1656	СВ	SER A 172	139.069	21.841	12.935	1.00 46.36
	ATOM	1657	OG	SER A 172	140.094	22.685	13.438	1.00 49.96
45	ATOM	1659	С	SER A 172	137.230	23.474	13.454	1.00 44.96
45	MOTA	1660	0	SER A 172	136.204	23.069	14.005	1.00 47.71
	MOTA	1661	N	GLN A 173	137.811	24.635	13.759	1.00 43.91
50	ATOM	1663	CA	GLN A 173	137.276	25.541	14.776	1.00 40.36
	ATOM	1664	СВ	GLN A 173	138.214	26.730	14.990	1.00 41.61
55	ATOM	1665	CG	GLN A 173	139.444	26.421	15.816	1.00 43.62
33	MOTA	1666	CD	GLN A 173	139.095	25.943	17.205	1.00 46.78
	MOTA	1667	OE1	GLN A 173	138.657	26.722	18.052	1.00 50.29
60	ATOM	1668	NE2	GLN A 173	139.279	24.654	17.448	1.00 48.32
	ATOM	1671	С	GLN A 173	135.893	26.047	14.386	1.00 38.81
65	ATOM	1672	0	GLN A 173	134.990	26.098	15.218	1.00 38.42
65	ATOM	1673	N	VAL A 174	135.734	26.423	13.120	1.00 36.59

	MOTA	1675	CA	VAL A 174	134.456	26.912	12.621	1.00 34.47
	ATOM	1676	СВ	VAL A 174	134.559	27.328	11.147	1.00 32.46
5	MOTA	1677	CG1	VAL A 174	133.183	27.559	10.564	1.00 32.42
	ATOM	1678	CG2	VAL A 174	135.380	28.601	11.037	1.00 31.13
10	ATOM	1679	С	VAL A 174	133.363	25.861	12.810	1.00 34.32
10	ATOM	1680	0	VAL A 174	132.350	26.130	13.445	1.00 36.14
	ATOM	1681	N	CYS A 175	133.586	24.652	12.308	1.00 35.02
15	ATOM	1683	CA	CYS A 175	132.601	23.591	12.460	1.00 35.40
	ATOM	1684	С	CYS A 175	132.440	23.152	13.906	1.00 36.62
20	MOTA	1685	0	CYS A 175	131.405	22.597	14.284	1.00 37.92
20	MOTA	1686	CB	CYS A 175	132.925	22.396	11.573	1.00 33.94
	ATOM	1687	SG	CYS A 175	132.302	22.595	9.876	1.00 42.37
25	ATOM	1688	N	GLU A 176	133.459	23.386	14.720	1.00 38.42
	MOTA	1690	CA	GLU A 176	133.365	23.034	16.123	1.00 40.14
30	MOTA	1691	CB	GLU A 176	134.739	23.093	16.793	1.00 43.35
50	MOTA	1692	CG	GLU A 176	134.747	22.502	18.193	1.00 50.93
	MOTA	1693	CD	GLU A 176	136.132	22.385	18.783	1.00 53.93
35	MOTA	1694	OE1	GLU A 176	136.867	21.459	18.382	1.00 58.04
	MOTA	1695	OE2	GLU A 176	136.481	23.207	19.658	1.00 57.02
40	MOTA	1696	C	GLU A 176	132.386	24.024	16.772	1.00 40.43
	MOTA	1697	0	GLU A 176	131.498	23.627	17.528	1.00 41.95
	MOTA	1698	N	LYS A 177	132.507	25.301	16.419	1.00 39.36
45	MOTA	1700	CA	LYS A 177	131.625	26.332	16.956	1.00 38.66
	ATOM	1701	CB	LYS A 177	132.074	27.729	16.510	1.00 38.94
50	MOTA	1702	CG	LYS A 177	133.403	28.194	17.089	1.00 40.75
	MOTA	1703	CD	LYS A 177	133.336	28.424	18.592	1.00 40.88
	MOTA	1704	CE	LYS A 177	134.694	28.852	19.152	1.00 40.13
55	ATOM	1705	NZ	LYS A 177	134.688	28.995	20.637	1.00 40.04
	ATOM	1709	С	LYS A 177	130.190	26.099	16.503	1.00 37.24
60	MOTA	1710	0	LYS A 177	129.277	26.078	17.318	1.00 39.39
	ATOM	1711	N	VAL A 178	130.002	25.901	15.204	1.00 35.46
	MOTA	1713	CA	VAL A 178	128.676	25.676	14.641	1.00 33.61
65	ATOM	1714	СВ	VAL A 178	128.747	25.442	13.124	1.00 31.21
	MOTA	1715	CG1	VAL A 178	127.365	25.166	12.568	1.00 32.70

	MOTA	1716	CG2	VAL A 178	129.346	26.652	12.446	1.00 31.16
_	ATOM	1717	С	VAL A 178	127.960	24.505	15.299	1.00 34.02
5	ATOM	1718	0	VAL A 178	126.844	24.656	15.784	1.00 36.09
	ATOM	1719	N	THR A 179	128.611	23.349	15.338	1.00 34.04
10	ATOM	1721	CA	THR A 179	128.018	22.167	15.947	1.00 35.60
	ATOM	1722	СВ	THR A 179	128.977	20.968	15.865	1.00 32.77
	ATOM	1723	OG1	THR A 179	129.372	20.775	14.505	1.00 32.73
15	ATOM	1725	CG2	THR A 179	128.296	19.706	16.330	1.00 35.18
	ATOM	1726	С	THR A 179	127.634	22.428	17.408	1.00 39.92
20	ATOM	1727	0	THR A 179	126.554	22.037	17.858	1.00 42.33
	ATOM	1728	N	ASN A 180	128.497	23.131	18.132	1.00 43.69
0.5	ATOM	1730	CA	ASN À 180	128.244	23.435	19.537	1.00 44.92
25	ATOM	1731	СВ	ASN A 180	129.496	23.991	20.213	1.00 49.20
	MOTA	1732	CG	ASN A 180	130.522	22.920	20.532	1.00 53.34
30	MOTA	1733	OD1	ASN A 180	131.550	23.213	21.140	1.00 58.92
	MOTA	1734	ND2	ASN A 180	130.258	21.680	20.126	1.00 53.15
25	MOTA	1737	С	ASN A 180	127.101	24.406	19.738	1.00 44.35
35	MOTA	1738	0	ASN A 180	126.252	24.194	20.593	1.00 46.90
	ATOM	1739	N	SER A 181	127.095	25.487	18.972	1.00 42.77
40	ATOM	1741	CA	SER A 181	126.047	26.481	19.084	1.00 42.55
	MOTA	1742	СВ	SER A 181	126.274	27.605	18.076	1.00 46.25
45	MOTA	1743	OG	SER A 181	127.508	28.261	18.319	1.00 51.45
43	ATOM	1745	С	SER A 181	124.686	25.841	18.869	1.00 41.46
	MOTA	1746	0	SER A 181	123.720	26.203	19.528	1.00 43.31
50	ATOM	1747	N	VAL A 182	124.610	24.870	17.969	1.00 41.03
	ATOM	1749	CA	VAL A 182	123.346	24.195	17.712	1.00 40.50
55	MOTA	1750	СВ	VAL A 182	123.468	23.177	16.565	1.00 40.39
J	MOTA	1751	CG1	VAL A 182	122.133	22.489	16.333	1.00 41.54
	MOTA	1752	CG2	VAL A 182	123.918	23.874	15.295	1.00 39.67
60	MOTA	1753	С	VAL A 182	122.865	23.481	18.970	1.00 41.04
	MOTA	1754	0	VAL A 182	121.831	23.836	19.536	1.00 41.32
C E	ATOM	1755	N	SER A 183	123.647	22.510	19.429	1.00 42.02
65	MOTA	1757	CA	SER A 183	123.311	21.735	20.616	1.00 41.82

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	MOTA	1758	CB	SER A	183	124.339	20.620	20.828	1.00 42.01
	MOTA	1759	OG	SER A	183	124.180	19.588	19.867	1.00 47.44
5	ATOM	1761	С	SER A	183	123.187	22.553	21.895	1.00 41.66
	MOTA	1762	0	SER A	183	122.251	22.358	22.673	1.00 43.45
10	ATOM-	1763	N	SER A	184	124.120	23.472	22.102	1.00 40.24
10	MOTA	1765	CA	SER A	184	124.143	24.293	23.305	1.00 40.26
	MOTA	1766	CB	SER A	184	125.590	24.639	23.670	1.00 42.61
15	MOTA	1767	OG	SER A	184	126.340	23.468	23.947	1.00 47.85
	MOTA	1769	С	SER A	184	123.307	25.567	23.312	1.00 39.22
20	MOTA	1770	0	SER A	184	122.991	26.088	24.382	1.00 40.56
20	MOTA	1771	N	GLU A	185	122.942	26.078	22.145	1.00 37.87
	MOTA	1773	CA	GLU A	185	122.169	27.313	22.098	1.00 37.71
25	MOTA	1774	СВ	GLU A	185	123.060	28.485	21.683	1.00 40.53
	ATOM	1775	CG	GLU A	185	124.010	28.960	22.768	1.00 45.79
30	MOTA	1776	CD	GLU A	185	124.987	30.024	22.296	1.00 50.89
50	MOTA	1777	OE1	GLU A	185	124.887	30.490	21.138	1.00 55.20
	ATOM	1778	OE2	GLU A	185	125.872	30.390	23.097	1.00 55.19
35	MOTA	1779	С	GLU A	185	120.934	27.277	21.225	1.00 35.83
	ATOM	1780	0	GLU A	185	119.921	27.861	21.584	1.00 38.77
40	ATOM	1781	N	LEU A	186	121.008	26.605	20.082	1.00 34.44
	ATOM	1783	CA	LEU A	186 .	119.864	26.533	19.187	1.00 33.39
	MOTA	1784	CB	LEU A	186	120.269	25.978	17.818	1.00 30.92
45	ATOM	1785	CG	LEU A	186	119.709	26.645	16.553	1.00 26.02
	ATOM	1786	CD1	LEU A	186	119.228	25.581	15.601	1.00 23.97
50	ATOM	1787	CD2	LEU A	186	118.581	27.619		1.00 25.38
	ATOM	1788	С	LEU A	186	118.794	25.652	19.802	1.00 34.20
	ATOM	1789	0	LEU A	186	117.655			1.00 35.35
55	ATOM	1790	N	GLN A	187	119.168	24.433	20.181	1.00 35.06
	ATOM	1792	CA	GLN A	187	118.218	23.500	20.778	1.00 37.32
60	MOTA	1793	CB		187	118.856	22.130	21.031	1.00 38.08
	ATOM	1794	CG		187	117.859	21.104		1.00 39.72
<i>-</i> -	MOTA	1795	CD		187	118.388			1.00 39.64
65	MOTA	1796			187	118.203			1.00 42.24
	MOTA	1797	NE2	GLN A	187	119.031	19.272	22.585	1.00 43.70

	MOTA	1800	C	GLN A 18	37 117.54	8 24.041	22.041	1.00	36.63
_	ATOM	1801	0	GLN A 18	37 116.32	5 24.022	22.138	1.00	40.27
5	MOTA	1802	N	PRO A 18	38 118.33	3 24.493	23.038	1.00	36.32
	MOTA	1803	CD	PRO A 18	38 119.78	2 24.293	23.223	1.00	38.65
10	MOTA	1804	CA	PRO A 18	38 117.74	8 25.033	24.268	1.00	35.72
	MOTA	1805	СВ	PRO A 18	38 118.97	7 25.432	25.075	1.00	34.46
1.5	ATOM	1806	CG	PRO A 18	38 119.93	3 24.364	24.730	1.00	35.29
15	MOTA	1807	С	PRO A 18	116.82	9 26.231	24.030	1.00	36.20
	MOTA	1808	0	PRO A 18	88 115.93	3 26.490	24.835	1.00	38.62
20	MOTA	1809	N	TYR A 18	89 117.06	2 26.983	22.957	1.00	35.08
	MOTA	1811	CA	TYR A 18	89 116.20	3 28.122	22.667	1.00	33.10
25	MOTA	1812	CB	TYR A 18	89 116.79	9 29.059	21.625	1.00	30.49
23	ATOM	1813	CG	TYR A 18	89 115.76	7 30.037	21.113	1.00	29.52
	ATOM	1814	CD1	TYR A 18	89 115.22	2 30.996	21.958	1.00	28.37
30	MOTA	1815	CE1	TYR A 1	89 114.21	1 31.842	21.523	1.00	27.60
	ATOM	1816	CD2	TYR A 1	89 115.27	4 29.950	19.808	1.00	28.40
35	ATOM	1817	CE2	TYR A 1	89 114.26	1 30.792	19.366	1.00	26.35
55	ATOM	1818	cz	TYR A 1	89 113.73	7 31.732	20.231	1.00	25.96
	ATOM	1819	OH	TYR A 1	89 112.73	5 32.568	19.815	1.00	27.51
40	MOTA	1821	С	TYR A 1	89 114.84	27.660	22.186	1.00	34.73
	MOTA	1822	0	TYR A 1	89 113.82	28.113	22.694	1.00	38.35
45	MOTA	1823	N	PHE A 1	90 114.81	1 26.771	21.197	1.00	36.21
	MOTA	1825	CA	PHE A 1	90 113.53	8 26.288	20.684	1.00	37.47
	MOTA	1826	CB	PHE A 1	90 113.69	25.649	19.302	1.00	38.76
50	ATOM	1827	CG	PHE A 1	90 113.52	22 26.633	18.177	1.00	41.54
	ATOM	1828	CD1	PHE A 1	.90 112.27	73 27.194	17.920	1.00	40.00
55	ATOM	1829	CD2	PHE A 1	.90 114.61	10 27.042	17.406	1.00	43.58
	MOTA	1830	CE1	PHE A 1	.90 112.10	9 28.146	16.919	1.00	39.03
	MOTA	1831	CE2	PHE A 1	.90 114.45	27.999	16.398	1.00	40.39
60	ATOM	1832	CZ	PHE A 1	.90 113.20	01 28.550	16.159	1.00	39.28
	ATOM	1833	С	PHE A 1	.90 112.76	68 25.408	21.662	1.00	37.92
65	MOTA	1834	0	PHE A 1	190 111.6	55 24.960	21.380	1.00	38.40
J.	ATOM	1835	N	GLN A 1	113.3	51 25.193	22.835	1.00	36.94

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	ATOM	1837	CA	GLN A	191	112.685	24.425	23.869	1.00	36.24-
	ATOM	1838	СВ	GLN A	191	113.637	23.426	24.529	1.00	36.06
5	ATOM	1839	CG	GLN A	191	113.987	22.259	23.609	1.00	37.66
	ATOM	1840	CD	GLN A	191	114.786	21.170	24.287	1.00	36.77
10	MOTA	1841	OE1	GLN A	191	115.365	21.379	25.349	1.00	38.68
10	MOTA	1842	NE2	GLN A	191	114.828	19.997	23.669	1.00	35.78
	ATOM	1845	С	GLN A	191	112.042	25.369	24.881	1.00	37.61
15	ATOM	1846	0	GLN A	191	111.621	24.945	25.956	1.00	38.67
	MOTA	1847	N	THR A	192	111.992	26.659	24.541	1.00	38.44
	ATOM	1849	CA	THR A	192	111.338	27.650	25.392	1.00	37.16
20	ATOM	1850	CB	THR A	192	111.946	29.055	25.263	1.00	36.82
	MOTA	1851	OG1	THR A	192	111.937	29.461	23.890	1.00	37.79
25	MOTA	1853	CG2	THR A	192	113.357	29.083	25.802	1.00	36.99
	MOTA	1854	C	THR A	192	109.894	27.702	24.913	1.00	37.14
30	ATOM	1855	0	THR A	192	109.082	28.485	25.403	1.00	38.53
50	ATOM	1856	N	LEU A	193	109.604	26.884	23.906	1.00	38.31
	ATOM	1858	CA	LEU A	193	108.273	26.765	23.339	1.00	40.73
35	ATOM	1859	CB	LEU A	193	108.325	25.772	22.174	1.00	42.93
	ATOM	1860	CG	LEU A	193	107.191	25.727	21.151	1.00	44.96
40	MOTA	1861	CD1	LEU A	193	107.000	27.100	20.518	1.00	45.48
	MOTA	1862	CD2	LEU A	193	107.531	24.696	20.085	1.00	44.59
	MOTA	1863	С	LEU A	193	107.403	26.218	24.472	1.00	40.91
45	ATOM	1864	0	LEU A	193	107.788	25.265	25.150	1.00	41.30
	MOTA	1865	N	PRO A	194	106.230	26.824	24.707	1.00	41.79
50	MOTA	1866	CD	PRO A	194	105.632	27.950	23.974	1.00	41.39
	MOTA	1867	CA	PRO A	194	105.334	26.373	25.777	1.00	41.80
	MOTA	1868	CB	PRO A		104.217	27.420	25.754		42.00
55	MOTA	1869	CG	PRO A	194	104.823	28.603	25.045	1.00	42.94
	MOTA	1870	С	PRO A	194	104.755	24.985	25.522	1.00	41.75
60	MOTA	1871	0	PRO A	194	104.174	24.734	24.469		44.72
	ATOM	1872	N	VAL A		104.930	24.080	26.476		39.33
_	ATOM	1874	CA	VAL A		104.383	22.738	26.338		37.64
65	ATOM	1875	CB	VAL A		105.403	21.659	26.756		37.05
	MOTA	1876	CG1	VAL A	195	104.744	20.295	26.840	1.00	34.56

	MOTA	1877	CG2	VAL A	195	106.515	21.607	25.747	1.00 36.43
E	ATOM	1878	С	VAL A	A 195	103.118	22.644	27.183	1.00 38.20
5	MOTA	1879	0	VAL A	195	102.062	22.245	26.694	1.00 40.04
	MOTA	1880	N	MET A	A 196	103.234	23.007	28.454	1.00 38.05
10	MOTA	1882	CA	MET A	196	102.103	22.987	29.370	1.00 37.71
	MOTA	1883	СВ	MET A	196	102.414	22.128	30.592	1.00 40.52
15	MOTA	1884	CG	MET A	196	102.756	20.681	30.283	1.00 43.85
15	MOTA	1885	SD	MET A	196	101.356	19.738	29.695	1.00 42.31
	MOTA	1886	CE	MET A	A 196	100.644	19.226	31.232	1.00 41.10
20	MOTA	1887	С	MET A	A 196	101.924	24.428	29.796	1.00 37.52
	MOTA	1888	0	MET A	A 196	102.521	24.877	30.771	1.00 40.72
25	ATOM	1889	N	THR A	A 197	101.134	25.167	29.039	1.00 36.00
25	ATOM	1891	CA	THR A	A 197	100.914	26.565	29.335	1.00 34.89
	MOTA	1892	СВ	THR A	A 197	100.650	27.324	28.036	1.00 37.85
30	MOTA	1893	OG1	THR A	A 197	101.329	26.660	26.963	1.00 41.31
	ATOM	1895	CG2	THR A	A 197	101.177	28.733	28.131	1.00 39.84
35	ATOM	1896	С	THR A	A 197	99.751	26.772	30.302	1.00 32.85
33	ATOM	1897	0	THR A	A 197	98.594	26.571	29.939	1.00 33.57
	MOTA	1898	Ŋ	LYS A	A 198	100.061	27.122	31.545	1.00 29.63
40	MOTA	1900	CA	LYS A	A 198	99.036	27.372	32.551	1.00 27.13
	ATOM	1901	СВ	LYS A	A 198	99.630	27.266	33.952	1.00 26.01
45	ATOM	1902	CG	LYS A	A 198	98.609	27.379	35.047	1.00 26.97
	ATOM	1903	CD	LYS 2	A 198	99.269	27.457	36.389	1.00 31.48
	MOTA	1904	CE	LYS 2	A 198	98.232	27.489	37.482	1.00 33.31
50	MOTA	1905	NZ	LYS 2	A 198	97.393	26.263	37.448	1.00 38.94
	MOTA	1909	С	LYS 2	A 198	98.555	28.789	32.307	1.00 26.60
55	MOTA	1910	0	LYS 2	A 198	99.365	29.706	32.264	1.00 27.53
	MOTA	1911	N	ILE A	A 199	97.248	28.975	32.154	1.00 26.62
	MOTA	1913	CA	ILE 2	A 199	96.711	30.298	31.878	1.00 23.94
60	MOTA	1914	CB	ILE	A 199	95.817	30.283	30.613	1.00 25.54
	ATOM ·	1915	CG2	ILE A	A 199	96.561	29.614	29.477	1.00 26.32
65	MOTA	1916	CG1	ILE	A 199	94.543	29.487	30.843	1.00 23.67
	MOTA	1917	CD1	ILE	A 199	93.795	29.189	29.564	1.00 22.95

	ATOM	1918	С	ILE A 199	95.992	30.954	33.041	1.00 25.86
	MOTA	1919	0	ILE A 199	95.827	32.169	33.061	1.00 28.61
5	MOTA	1920	N	ASP A 200	95.561	30.148	34.007	1.00 27.99
	MOTA	1922	CA	ASP A 200	94.878	30.651	35.197	1.00 26.92
10	ATOM	1923	СВ	ASP A 200	93.484	31.220	34.867	1.00 26.06
10	MOTA	1924	CG	ASP A 200	92.582	30.234	34.151	1.00 24.60
	MOTA	1925	OD1	ASP A 200	92.621	29.037	34.465	1.00 26.68
15	MOTA	1926	OD2	ASP A 200	91.813	30.664	33.271	1.00 28.99
	MOTA	1927	С	ASP A 200	94.805	29.589	36.289	1.00 27.44
20	MOTA	1928	0	ASP A 200	95.509	28.591	36.225	1.00 29.73
20	MOTA	1929	N	SER A 201	93.966	29.808	37.294	1.00 28.30
	MOTA	1931	CA	SER A 201	93.825	28.859	38.397	1.00 30.40
25	MOTA	1932	CB	SER A 201	93.032	29.507	39.531	1.00 31.85
	ATOM	1933	OG	SER A 201	91.882	30.170	39.026	1.00 39.39
30	ATOM	1935	С	SER A 201	93.137	27.562	38.000	1.00 29.79
30	ATOM	1936	0	SER A 201	93.230	26.557	38.695	1.00 31.67
	ATOM	1937	N	VAL A 202	92.430	27.604	36.884	1.00 30.29
35	MOTA	1939	CA	VAL A 202	91.683	26.457	36.409	1.00 29.89
	MOTA	1940	CB	VAL A 202	90.284	26.911	35.932	1.00 28.59
40	ATOM	1941	CG1	VAL A 202	89.426	25.723	35.568	1.00 30.45
40	ATOM	1942	CG2	VAL A 202	89.613	27.740	36.998	1.00 28.22
	MOTA	1943	С	VAL A 202	92.346	25.667	35.282	1.00 31.71
45	ATOM	1944	0	VAL A 202	92.752	24.520	35.470	1.00 33.93
	ATOM	1945	N	ALA A 203	92.463	26.291	34.115	1.00 31.10
50	MOTA	1947	CA	ALA A 203	92.997	25.621	32.944	1.00 29.58
	MOTA	1948	CB	ALA A 203	92.039	25.821	31.767	1.00 29.01
	MOTA	1949	C	ALA A 203	94.416	25.908	32.497	1.00 28.91
55	MOTA	1950	0	ALA A 203	95.096	26.807	33.001	1.00 27.86
	ATOM	1951	N	GLY A 204	94.822	25.124	31.503	1.00 26.23
60	MOTA	1953	CA	GLY A 204	96.124	25.227	30.894	1.00 22.94
	MOTA	1954	С	GLY A 204	95.936	24.686	29.495	1.00 22.27
	ATOM	1955	0	GLY A 204	94.900	24.098	29.206	1.00 24.27
65	MOTA	1956	N	ILE A 205	96.909	24.893	28.619	1.00 22.81
	MOTA	1958	CA	ILE A 205	96.817	24.402	27.250	1.00 21.62

	ATOM	1959	CB	ILE .	A	205	96.868	25.542	26.222	1.00	17.56
-	ATOM	1960	CG2	ILE .	A	205	96.458	25.020	24.870	1.00	17.14
5	ATOM	1961	CG1	ILE .	A	205	95.928	26.672	26.627	1.00	16.69
	ATOM	1962	CD1	ILE .	A	205	96.021	27.878	25.753	1.00	10.87
10	ATOM	1963	C	ILE .	A	205	98.010	23.492	27.008	1.00	24.34
	MOTA	1964	0	ILE .	A	205	99.120	23.784	27.452	1.00	26.93
15	ATOM	1965	N	ASN .	Α	206	97.770	22.374	26.333	1.00	24.44
15	ATOM	1967	CA	ASN .	A	206	98.817	21.414	26.039	1.00	20.90
	ATOM	1968	СВ	ASN	A	206	98.280	19.989	26.217	1.00	21.75
20	ATOM	1969	CG	ASN	A	206	99.359	18.921	26.099	1.00	22.44
	ATOM	1970	OD1	ASN	A	206	99.077	17.727	26.218	1.00	26.27
25	ATOM	1971	ND2	ASN	A	206	100.593	19.337	25.878	1.00	24.68
	ATOM	1974	С	ASN	A	206	99.285	21.629	24.614	1.00	22.48
	ATOM	1975	0	ASN	A	206	98.579	21.296	23.664	1.00	23.50
30	ATOM	1976	N	TYR	A	207	100.458	22.239	24.475	1.00	23.09
	ATOM	1978	CA	TYR	A	207	101.065	22.517	23.179	1.00	21.41
35	ATOM	1979	CB	TYR	A	207	101.647	23.927	23.165	1.00	17.69
55	MOTA	1980	CG	TYR	A	207	100.625	25.008	22.986	1.00	16.56
	MOTA	1981	CD1	TYR	A	207	99.703	24.943	21.961	1.00	19.32
40	MOTA	1982	CE1	TYR	A	207	98.795	25.963	21.757	1.00	20.24
	MOTA	1983	CD2	TYR	A	207	100.611	26.122	23.810	1.00	18.10
45	MOTA	1984	CE2	TYR	A	207	99.711	27.144	23.613	1.00	18.49
	MOTA	1985	CZ	TYR	A	207	98.809	27.060	22.581	1.00	18.16
	MOTA	1986	OH	TYR	A	207	97.939	28.089	22.332	1.00	21.72
50	MOTA	1988	С	TYR	A	207	102.181	21.532	22.868	1.00	24.14
	MOTA	1989	0	TYR	A	207	103.111	21.851	22.134	1.00	25.16
55	MOTA	1990	N	GLY	A	208	102.083	20.328	23.418	1.00	28.50
	ATOM	1992	CA	GLY	A	208	103.112	19.330	23.194	1.00	29.80
	ATOM	1993	С	GLY	A	208	103.130	18.752	21.795	1.00	32.56
60	ATOM	1994		GLY	A	208	102.067	18.467	21.231	1.00	33.45
	ATOM	1995	N	LEU	A	209	104.332	18.598	21.235	1.00	33.15
65	ATOM	1997	CA	LEU	A	209	104.512	18.038	19.899	1.00	31.79
	ATOM	1998	CB	LEU	A	209	105.985	18.081	19.484	1.00	29.49

	MOTA	1999	CG	LEU A	209	106.491	19.337	18.773	1.00	28.06
	MOTA	2000	CD1	LEU A	209	106.302	20.558	19.641	1.00	24.35
5	ATOM	2001	CD2	LEU A	209	107.953	19.158	18.416	1.00	24.06
	MOTA	2002	C	LEU A	209	104.052	16.599	19.941	1.00	31.52
10	ATOM	2003	0	LEU A	209	104.416	15.863	20.854	1.00	33.82
10	MOTA	2004	N	VAL A	210	103.254	16.199	18.961	1.00	29.87
	ATOM	2006	CA	VAL A	210	102.752	14.836	18.915	1.00	29.99
15	MOTA	2007	СВ	VAL A	210	101.226	14.817	18.732	1.00	30.37
	MOTA	2008	CG1	VAL A	210	100.559	15.363	19.972	1.00	31.24
20	MOTA	2009	CG2	VAL A	210	100.824	15.642	17.522	1.00	30.83
20	MOTA	2010	С	VAL A	210	103.430	13.992	17.837	1.00	30.78
	MOTA	2011	0	VAL A	210	103.038	12.854	17.578	1.00	32.27
25 🦏	MOTA	2012	N	ALA A	211	104.460	14.552	17.221	1.00	30.97
	MOTA	2014	CA	ALA A	211	105.199	13.857	16.178	1.00	32.26
30	MOTA	2015	СВ	ALA A	211	104.324	13.661	14.941	1.00	34.28
50	MOTA	2016	С	ALA A	211	106.417	14.692	15.839	1.00	31.64
	MOTA	2017	0	ALA A	211	106.463	15.884	16.145	1.00	32.86
35	MOTA	2018	N	PRO A	212	107.437	14.077	15.232	1.00	30.38
	ATOM	2019	CD	PRO A	212	107.550	12.667	14.820	1.00	28.87
40	MOTA	2020	CA	PRO A	212	108.642	14.824	14.876	1.00	30.59
	MOTA	2021	CB	PRO A	212	109.576	13.724	14.384	1.00	30.90
	MOTA	2022	CG	PRO A	212	108.628	12.731	13.785	1.00	31.64
45	MOTA	2023	С	PRO A	212	108.345	15.824	13.774	1.00	31.31
	ATOM	2024	0	PRO A	212	107.483	15.574	12.938	1.00	33.01
50	ATOM	2025	N	PRO A	213	109.005	16.996	13.802	1.00	30.25
	MOTA	2026	CD	PRO A	213	109.882	17.457	14.892	1.00	28.21
	ATOM	2027	CA	PRO A	213	108.832	18.056	12.803	1.00	29.74
55	MOTA	2028	CB	PRO A	213	109.889	19.076	13.220	1.00	29.54
	MOTA	2029	CG	PRO A	213	109.896	18.958	14.690	1.00	26.77
60	MOTA	2030	С	PRO A	213	109.115	17.526	11.398	1.00	30.72
	MOTA	2031	0	PRO A	213	110.187	16.975	11.140	1.00	33.55
	ATOM	2032	N	ALA A	214	108.157	17.700	10.495	1.00	30.16
65	ATOM	2034	CA	ALA A	214	108.296	17.231	9.123	1.00	29.31
	MOTA	2035	CB	ALA A	214	106.964	16.701	8.609	1.00	27.36

	ATOM	2036	С	ALA A 214	108.771	18.336	8.212	1.00 30.52
£	ATOM	2037	0	ALA A 214	108.229	19.440	8.235	1.00 32.72
5	ATOM	2038	N	THR A 215	109.781	18.052	7.405	1.00 30.13
	ATOM	2040	CA	THR A 215	110.268	19.047	6.472	1.00 31.47
10	ATOM	2041	СВ	THR A 215	111.805	19.044	6.366	1.00 32.14
	ATOM	2042	OG1	THR A 215	112.381	19.228	7.665	1.00 36.33
	ATOM	2044	CG2	THR A 215	112.268	20.177	5.474	1.00 33.34
15	ATOM	2045	С	THR A 215	109.656	18.698	5.127	1.00 32.34
	ATOM	2046	0	THR A 215	109.642	17.533	4.729	1.00 32.18
20	ATOM	2047	N	THR A 216	109.103	19.692	4.451	1.00 34.34
	MOTA	2049	CA	THR A 216	108.489	19.463	3.154	1.00 36.27
25	MOTA	2050	СВ	THR A 216	106.996	19.862	3.167	1.00 36.83
25	MOTA	2051	OG1	THR A 216	106.872	21.255	3.477	1.00 37.89
	MOTA	2053	CG2	THR A 216	106.241	19.055	4.216	1.00 36.73
30	MOTA	2054	С	THR A 216	109.247	20.254	2.092	1.00 37.93
	MOTA	2055	0	THR A 216	110.390	20.642	2.307	1.00 36.96
35	ATOM	2056	N	ALA A 217	108.610	20.483	0.948	1.00 40.73
33	MOTA	2058	CA	ALA A 217	109.228	21.215	-0.151	1.00 41.25
	MOTA	2059	СВ	ALA A 217	108.452	20.967	-1.437	1.00 42.73
40	ATOM	2060	С	ALA A 217	109.302	22.704	0.125	1.00 40.38
	MOTA	2061	0	ALA A 217	110.218	23.379	-0.337	1.00 40.36
45	MOTA	2062	N	GLU A 218	108.319	23.215	0.860	1.00 41.20
13	MOTA	2064	CA	GLU A 218	108.271	24.633	1.170	1.00 40.88
	MOTA	2065	CB	GLU A 218	107.443	25.372	0.113	1.00 46.00
50	MOTA	2066	CG	GLU A 218	108.304	26.058	-0.947	1.00 53.45
	MOTA	2067	CD	GLU A 218	107.610	26.213	-2.286	1.00 57.50
55	ATOM	2068	OEI	GLU A 218	106.543	25.588	-2.487	1.00 60.01
20	ATOM	2069	OE2	2 GLU A 218	108.149	26.945	-3.146	1.00 59.32
	ATOM	2070	С	GLU A 218	107.832	25.030	2.580	1.00 39.13
60	ATOM	2071	0	GLU A 218	107.635	26.216	2.839	1.00 40.54
	MOTA	2072	N	THR A 219	107.668	24.066	3.488	1.00 37.20
65	ATOM	2074	CA	THR A 219	107.295	24.388	4.875	1.00 35.55
55	MOTA	2075	CB	THR A 219	105.762	24.464	5.103	1.00 32.68

	ATOM	2076	OG1	THR A 219	105.144	23.249	4.674	1.00 35.48
	MOTA	2078	CG2	THR A 219	105.151	25.641	4.387	1.00 33.81
5	MOTA	2079	С	THR A 219	107.828	23.400	5.906	1.00 34.31
	ATOM	2080	0	THR A 219	108.013	22.217	5.614	1.00 34.57
10	ATOM	2081	N	LEU A 220	108.087	23.903	7.109	1.00 31.67
10	ATOM	2083	CA	LEU A 220	108.533	23.079	8.219	1.00 29.70
	ATOM	2084	СВ	LEU A 220	109.587	23.834	9.033	1.00 29.62
15	ATOM	2085	CG	LEU A 220	110.345	23.149	10.177	1.00 31.10
	ATOM	2086	CD1	LEU A 220	109.511	23.047	11.435	1.00 30.08
20	ATOM	2087	CD2	LEU A 220	110.794	21.791	9.727	1.00 30.78
20	ATOM	2088	С	LEU A 220	107.236	22.902	9.017	1.00 29.78
``	ATOM	2089	0	LEU A 220	106.693	23.878	9.545	1.00 30.49
25 `	ATOM	2090	N	ASP A 221	106.698	21.685	9.032	1.00 29.15
	MOTA	2092	CA	ASP A 221	105.441	21.397	9.728	1.00 30.48
30	ATOM	2093	СВ	ASP A 221	104.603	20.394	8.924	1.00 34.57
30	MOTA	2094	CG	ASP A 221	104.178	20.924	7.557	1.00 37.91
	MOTA	2095	OD1	ASP A 221	104.480	22.090	7.222	1.00 37.84
35	MOTA	2096	OD2	ASP A 221	103.534	20.158	6.811	1.00 42.30
	MOTA	2097	С	ASP A 221	105.607	20.865	11.143	1.00 30.64
40	MOTA	2098	0	ASP A 221	106.274	19.852	11.350	1.00 31.22
40	MOTA	2099	N	VAL A 222	104.950	21.519	12.100	1.00 29.81
	MOTA	2101	CA	VAL A 222	104.999	21.131	13.512	1.00 29.49
45	MOTA	2102	CB	VAL A 222	105.588	22.261	14.389	1.00 29.52
	MOTA	2103	CG1	VAL A 222	105.617	21.831	15.845	1.00 28.17
50	ATOM	2104	CG2	VAL A 222	106.988	22.632	13.914	1.00 26.55
	ATOM	2105	С	VAL A 222	103.586	20.791	14.016	1.00 29.65
	MOTA	2106	0	VAL A 222	102.686	21.620	13.958	1.00 31.27
55	MOTA	2107	N	GLN A 223	103.400	19.560	14.485	1.00 29.35
	MOTA	2109	CA	GLN A 223	102.114	19.081	14.978	1.00 25.88
60	ATOM	2110	СВ	GLN A 223	101.870	17.649	14.488	1.00 29.16
	MOTA	2111	CG	GLN A 223	101.731	17.505	12.969	1.00 30.27
	ATOM	2112	CD	GLN A 223	101.704	16.052	12.510	1.00 32.10
65	MOTA	2113	OE1	GLN A 223	100.832	15.275	12.897	1.00 31.46
	MOTA	2114	NE2	GLN A 223	102.668	15.681	11.686	1.00 37.75

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	ATOM	2117	С	GLN A 223	102.081	19.116	16.494	1.00 24.57
_	MOTA	2118	0	GLN A 223	102.910	18.490	17.147	1.00 23.51
5	MOTA	2119	N	MET A 224	101.112	19.840	17.042	1.00 22.94
	MOTA	2121	CA	MET A 224	100.959	19.984	18.480	1.00 22.16
10	MOTA	2122	СВ	MET A 224	101.064	21.459	18.866	1.00 23.89
	MOTA	2123	CG	MET A 224	102.353	22.116	18.423	1.00 22.07
1.5	MOTA	2124	SD	MET A 224	102.393	23.853	18.797	1.00 28.37
15	MOTA	2125	CE	MET A 224	104.122	24.130	18.771	1.00 25.60
	MOTA	2126	С	MET A 224	99.620	19.408	18.927	1.00 23.31
20	MOTA	2127	0	MET A 224	98.702	19.294	18.124	1.00 27.48 .
	MOTA	2128	N	LYS A 225	99.519	19.024	20.197	1.00 25.28
25	MOTA	2130	CA	LYS A 225	98.292	18.440	20.735	1.00 27.30
23	MOTA	2131	СВ	LYS A 225	98.522	17.889	22.140	1.00 28.45
	MOTA	2132	CG	LYS A 225	97.613	16.724	22.452	1.00 30.88
30	MOTA	2133	CD	LYS A 225	97.912	16.108	23.793	1.00 33.18
	MOTA	2134	CE	LYS A 225	97.153	14.802	23.938	1.00 38.68
35	ATOM	2135	NZ	LYS A 225	95.741	14.928	23.460	1.00 43.39
55	ATOM	2139	C	LYS A 225	97.135	19.434	20.726	1.00 27.30
	MOTA	2140	0	LYS A 225	96.052	19.123	20.238	1.00 29.51
40	ATOM	2141	N,	GLY A 226 .	97.364	20.617	21.286	1.00 26.18
	MOTA	2143	CA	GLY A 226	96.360	21.665	21.285	1.00 21.95
45	MOTA	2144	С	GLY A 226	95.015	21.379	21.903	1.00 23.48
	MOTA	2145	0	GLY A 226	93.983	21.420	21.238	1.00 23.78
	MOTA	2146	N	GLU A 227	95.021	21.131	23.201	1.00 26.02
50	MOTA	2148	CA	GLU A 227	93.792	20.882	23.915	1.00 26.24
	MOTA	2149	CB	GLU A 227	93.547	19.384	24.068	1.00 29.17
55	ATOM	2150	CG	GLU A 227	94.467	18.697	25.052	1.00 25.81
	MOTA	2151	CD	GLU A 227	93.967	17.330	25.456	1.00 29.15
	ATOM	2152	OE1	GLU A 227	93.343	16.654	24.614	1.00 33.43
60	ATOM	2153	OE2	2 GLU A 227	94.199	16.928	26.615	1.00 31.54
	ATOM	2154	С	GLU A 227	93.937	21.514	25.283	1.00 27.06
65	ATOM	2155	0	GLU A 227	95.006	21.447	25.892	1.00 26.79
	MOTA	2156	N	PHE A 228	92.904	22.219	25.722	1.00 26.37

	MOTA	2158	CA	PHE A	A	228	92.942	22.816	27.034	1.00	24.07
	ATOM	2159	CB	PHE A	A	228	91.785	23.792	27.242	1.00	24.55
5	ATOM	2160	CG	PHE 2	A	228	91.987	25.134	26.592	1.00	23.72
	ATOM	2161	CD1	PHE 2	Α	228	92.066	25.251	25.207	1.00	21.59
10	ATOM	2162	CD2	PHE 2	A	228	92.045	26.287	27.366	1.00	21.39
10	ATOM	2163	CE1	PHE 2	A	228	92.193	26.498	24.607	1.00	21.74
	ATOM	2164	CE2	PHE A	A	228	92.172	27.531	26.779	1.00	21.24
15	ATOM	2165	CZ	PHE 2	A	228	92.246	27.640	25.395	1.00	24.55
	MOTA	2166	c ·	PHE A	A	228	92.779	21.630	27.949	1.00	25.56
~20	ATOM	2167	0	PHE 2	A	228	92.069	20.680	27.624	1.00	26.15
20	ATOM	2168	N	TYR Z	A	229	93.479	21.655	29.067	1.00	29.13
	MOTA	2170	CA	TYR 2	A	229	93.400	20.573	30.025	1.00	34.87
25	MOTA	2171	СВ	TYR .	A	229	94.735	19.820	30.084	1.00	32.93
	MOTA	2172	CG	TYR .	A	229	95.870	20.654	30.639	1.00	33.77
30	MOTA	2173	CD1	TYR Z	A	229	96.034	20.816	32.015	1.00	34.08
30	ATOM	2174	CE1	TYR 2	A	229	97.021	21.637	32.530	1.00	33.96
	MOTA	2175	CD2	TYR 2	A	229	96.739	21.332	29.793	1.00	34.25
35	ATOM	2176	CE2	TYR .	A	229	97.735	22.152	30.302	1.00	34.53
	ATOM	2177	CZ	TYR .	A	229	97.863	22.302	31.670	1.00	33.54
40	ATOM	2178	ОН	TYR 2	A	229	98.819	23.138	32.185	1.00	38.50
	MOTA	2180	С	TYR .	A	229	93.097	21.196	31.380	1.00	39.53
	MOTA	2181	0	TYR .	A	229	93.192	22.414	31.545	1.00	39.55
45	MOTA	2182	N	SER .	A	230	92.738	20.355	32.341	1.00	46.28
	MOTA	2184	CA	SER .	A	230	92.453	20.796	33.699	1.00	53.45
50	MOTA	2185	CB	SER .	A	230	90.998	20.491	34.071	1.00	54.47
	MOTA	2186	OG	SER .	A	230	90.720	20.820	35.424	1.00	56.07
	MOTA	2188	С	SER	A	230	93.396	19.996	34.582	1.00	58.48
55	ATOM	2189	0	SER	A	230	93.659	18.826	34.301	1.00	58.09
	MOTA	2190	N	GLU	A	231	93.946	20.632	35.611	1.00	66.92
60	ATOM	2192	CA	GLU	A	231	94.860	19.940	36.517	1.00	74.85
	ATOM	2193	CB	GLU	A	231	95.976	20.878	36.995	1.00	77.02
.	ATOM	2194	CG	GLU			97.377	20.450	36.548		81.62
65	ATOM	2195	CD			231	98.476	21.372			84.67
	MOTA	2196	OE1	GLU	A	231	98.402	22.592	36.790	1.00	86.64

	ATOM	2197	OE2	GLU A 2	231	99.421	20.876	37.709	1.00	85.89
_	ATOM	2198	С	GLU A 2	231	94.119	19.328	37.706	1.00	78.17
5	ATOM	2199	0	GLU A 2	231	94.602	18.379	38.323	1.00	79.46
	ATOM	2200	N	ALA A 2	232	92.934	19.855	38.003	1.00	81.24
10	ATOM	2202	CA	ALA A 2	232	92.131	19.363	39.118	1.00	84.16
	ATOM	2203	СВ	ALA A 2	232	91.111	20.417	39.544	1.00	84.14
. ~	ATOM	2204	С	ALA A 2	232	91.426	18.053	38.777	1.00	86.04
15	ATOM	2205	0	ALA A 2	232	91.078	17.279	39.671	1.00	87.55
	MOTA	2206	N	ALA A 2	233	91.203	17.819	37.487	1.00	86.79
20	MOTA	2208	CA	ALA A 2	233	90.534	16.608	37.023	1.00	88.81
	MOTA	2209	CB	ALA A 2	233	89.052	16.659	37.381	1.00	89.87
25	MOTA	2210	С	ALA A 2	233	90.702	16.464	35.516	1.00	90.03
25	ATOM	2211	0	ALA A 2	233	90.288	17.341	34.756	1.00	90.77
	ATOM	2212	N	ALA A	234	91.309	15.361	35.090	1.00	91.16
30	MOTA	2214	CA	ALA A	234	91.536	15.106	33.670	1.00	91.66
	MOTA	2215	CB	ALA A	234	92.810	14.288	33.470	1.00	92.37
35	MOTA	2216	С	ALA A	234	90.354	14.394	33.034	1.00	91.31
33	MOTA	2217	0	ALA A	234	89.874	13.384	33.550	1.00	91.09
	MOTA	2218	N	ALA A	235	89.894	14.926	31.908	1.00	91.34
40	MOTA	2220	CA	ALA A	235	88.774	14.346	31.180	1.00	90.88
	MOTA	2221	CB	ALA A	235	87.457	14.975	31.627	1.00	91.57
45	MOTA	2222	С	ALA A	235	88.993	14.572	29.691	1.00	89.83
	MOTA	2223	0	ALA A	235	89.205	15.706	29.247	1.00	89.77
	ATOM	2224	N	PRO A	236	89.032	13.480	28.912	1.00	88.56
50	ATOM	2225	CD	PRO A	236	88.962	12.083	29.376	1.00	88.69
	MOTA	2226	CA	PRO A	236	89.232	13.539	27.463	1.00	85.91
55	ATOM	2227	CB	PRO A	236	89.511	12.080	27.104	1.00	86.56
	ATOM	2228	ÇG	PRO A	236	88.680	11.334	28.099	1.00	88.16
	MOTA	2229	С	PRO A	236	87.997	14.085	26.738	1.00	82.58
60	ATOM	2230	0	PRO A	236	86.861	13.877	27.172	1.00	81.75
	MOTA	2231	N	PRO A	237	88.217	14.829	25.644	1.00	79.77
65	MOTA	2232	CD	PRO A	237	89.551	15.264	25.202	1.00	80.44
	ATOM	2233	CA	PRO A	237	87.178	15.444	24.811	1.00	77.34

	ATOM	2234	CB	PRO P	237	87.985	16.332	23.863	1.00 7	7.69
	ATOM	2235	CG	PRO F	237	89.249	16.604	24.617	1.00 7	9.85
.5	ATOM	2236	С	PRO A	237	86.366	14.426	24.015	1.00 7	5.23
	ATOM	2237	0	PRO P	237	86.919	13.462	23.480	1.00 7	5.90
10	MOTA	2238	N	PRO A	238	85.042	14.634	23.918	1.00 7	2.71
10	ATOM	2239	CD	PRO A	238	84.282	15.606	24.724	1.00 7	2.84
	ATOM	2240	CA	PRO A	238	84.131	13.750	23.182	1.00 7	0.12
15	ATOM	2241	СВ	PRO A	238	82.774	14.086	23.793	1.00 7	1.89
	ATOM	2242	CG	PRO F	238	82.907	15.535	24.109	1.00 7	2.51
20	ATOM	2243	С	PRO A	238	84.146	14.004	21.672	1.00 6	7.49
20	ATOM	2244	0	PRO A	238	83.099	14.016	21.017	1.00 6	6.70
	ATOM	2245	N	PHE A	3 239	85.340	14.216	21.130	1.00 6	4.79
25	ATOM	2247	CA	PHE P	A 239	85.521	14.472	19.706	1.00 6	1.01
	ATOM	2248	СВ	PHE A	A 239	84.959	15.849	19.319	1.00 6	0.29
30	ATOM	2249	CG	PHE A	239	85.172	16.923	20.362	1.00 5	7.87
50	ATOM	2250	CD1	PHE A	A 239	86.307	17.721	20.339	1.00 5	6.42
	ATOM	2251	CD2	PHE A	A 239	84.216	17.159	21.343	1.00 5	5.75
35	ATOM	2252	CEl	PHE A	239	86.484	18.737	21.274	1.00 5	3.82
	ATOM	2253	CE2	PHE A	A 239	84.387	18.171	22.279	1.00 5	3.85
40	ATOM	2254	CZ	PHE A	A 239	85.522	18.960	22.243	1.00 5	2.72
	ATOM	2255	С	PHE A	A 239 '	86.989	14.355	19.307	1.00 5	8.61
	MOTA	2256	0	PHE A	A 239	87.873	14.266	20.164	1.00 5	8.11
45	ATOM	2257	N	ALA A	4 240	87.234	14.314	18.002	1.00 5	5.15
	ATOM	2259	CA	ALA A	4 240	88.586	14.203	17.472	1.00 5	1.42
50	ATOM	2260	CB	ALA A	A 240	88.716	12.936	16.631	1.00 5	1.91
	MOTA	2261	С	ALA I	A 240	88.884.	15.441	16.629	1.00 4	19.88
	MOTA	2262	0	ALA 1	A 240	87.972	16.034	16.048	1.00 4	8.56
55	MOTA	2263	N	PRO 2	A 241	90.155	15.885	16.604	1.00 4	18.54
	MOTA	2264	CD	PRO I	A 241	91.270	15.398	17.436	1.00 4	17.09
60	MOTA	2265	CA	PRO 2	A 241	90.567	17.064	15.830	1.00 4	16.53
	MOTA	2266	CB	PRO 2	A 241	92.043	17.218	16.205	1.00 4	14.66
	MOTA	2267	CG	PRO I	A 241	92.113	16.633	17.574	1.00 4	15.79
65	MOTA	2268	С	PRO 2	A 241	90.412	16.864	14.322	1.00 4	16.25
	MOTA	2269	0	PRO 2	A 241	90.536	15.743	13.819	1.00 4	17.76

	MOTA	2270	N	PRO A	242	90.118	17.947	13.582	1.00	45.32
_	MOTA	2271	CD	PRO A 2	242	89.818	19.314	14.050	1.00	44.47
5	MOTA	2272	CA	PRO A	242	89.959	17.844	12.130	1.00	44.33
	MOTA	2273	CB	PRO A	242	89.187	19.114	11.792	1.00	43.70
10	MOTA	2274	CG	PRO A	242	89.769	20.107	12.755	1.00	43.85
	MOTA	2275	С	PRO A	242	91.325	17.835	11.442	1.00	45.10
1.5	MOTA	2276	0	PRO A	242	92.331	18.261	12.019	1.00	44.03
15	ATOM	2277	N	VAL A	243	91.370	17.302	10.228	1.00	45.79
	MOTA	2279	CA	VAL A	243	92.615	17.279	9.485	1.00	47.91
20	MOTA	2280	СВ	VAL A	243	92.613	16.226	8.347	1.00	48.72
	MOTA	2281	CG1	VAL A	243	92.629	14.827	8.934	1.00	50.59
25	MOTA	2282	CG2	VAL A	243	91.403	16.401	7.446	1.00	51.20
25	MOTA	2283	C	VAL A	243	92.753	18.679	8.926	1.00	48.35
	MOTA	2284	0	VAL A	243	91.939	19.122	8.116	1.00	48.79
30	MOTA	2285	N	MET A	244	93.731	19.412	9.434	1.00	48.68
	ATOM	2287	CA	MET A	244	93.937	20.771	8.987	1.00	46.91
35	ATOM	2288	CB	MET A	244	94.406	21.638	10.147	1.00	42.52
33	ATOM	2289	CG	MET A	244	93.369	21.770	11.233	1.00	30.61
	ATOM	2290	SD	MET A	244	93.963	22.834	12.505	1.00	30.25
40	MOTA	2291	CE	MET A	244	94.948	21.710	13.447	1.00	25.62
	MOTA	2292	С	MET A	244	94.884	20.877	7.811	1.00	50.14
45	MOTA	2293	0	MET A	244	96.102	20.935	7.983	1.00	50.38
	MOTA	2294	N	GLU A	245	94.302	20.867	6.615	1.00	53.76
	MOTA	2296	CA	GLU A	245	95.046	20.985	5.367	1.00	58.60
50	MOTA	2297	СВ	GLU A	245	94.597	19.905	4.364	1.00	61.51
	MOTA	2298	CG	GLU A	245	93.085	19.861	4.078	1.00	67.59
55	MOTA	2299	CD	GLU A	245	92.692	18.811	3.041	1.00	69.79
	MOTA	2300	OE1	GLU A	245	92.943	19.024	1.827	1.00	72.17
	ATOM	2301	OE2	GLU A	245	92.104	17.778	3.439	1.00	68.95
60	ATOM	2302	С	GLU A	245	94.817	22.382	4.794	1.00	59.59
	ATOM	2303	0	GLU A	245	93.765	22.664	4.220	1.00	60.79
65	ATOM	2304	N	PHE A	246	95 . 773	23.278	5.010	1.00	61.44
03	ATOM	2306	CA	PHE A	246	95.637	24.635	4.510	1.00	63.27

	MOTA	2307	CB	PHE A 24	95.567	25.658	5.657	1.00	60.46°
	MOTA	2308	CG	PHE A 246	96.689	25.567	6.663	1.00	56.43
5	MOTA	2309	CD1	PHE A 246	97.850	26.319	6.502	1.00	54.28
	MOTA	2310	CD2	PHE A 240	96.532	24.825	7.834	1.00	53.72
10	ATOM	2311	CE1	PHE A 24	98.829	26.342	7.497	1.00	51.24
10	ATOM	2312	CE2	PHE A 240	97.506	24.841	8.834	1.00	51.21
	MOTA	2313	CZ	PHE A 24	98.653	25.602	8.667	1.00	48.89
15	MOTA	2314	С	PHE A 24	96.676	25.022	3.473	1.00	67.53
	MOTA	2315	0	PHE A 240	97.723	24.382	3.357	1.00	68.37
20	ATOM	2316	N	PRO A 24	96.375	26.050	2.664	1.00	71.27
20	ATOM	2317	CD	PRO A 24	95.153	26.874	2.673	1.00	72.37
	MOTA	2318	·CA	PRO A 24	7 97.297	26.511	1.624	1.00	74.18
25	ATOM	2319	CB	PRO A 24	7 96.504	27.627	0.938	1.00	74.64
	MOTA	2320	CG	PRO A 24	7 95.626	28.147	2.038	1.00	73.71
30	MOTA	2321	c	PRO A 24	7 98.620	27.016	2.180	1.00	76.26
50	MOTA	2322	0	PRO A 24	7 98.661	28.002	2.920	1.00	76.91
	MOTA	2323	N	ALA A 24	99.691	26.299	1.858	1.00	78.37
35	MOTA	2325	CA	ALA A 24	101.027	26.675	2.298	1.00	80.34
	MOTA	2326	СВ	ALA A 24	102.025	25.578	1.963	1.00	80.57
40	MOTA	2327	С	ALA A 24	101.385	27.950	1.556	1.00	81.06
•	MOTA	2328	0	ALA A 24	8 101.463	27.962	0.325	1.00	81.92
	MOTA	2329	N	ALA A 24	9 101.564	29.030	2.300	1.00	80.72
45	MOTA	2331	CA	ALA A 24	9 101.902	30.303	1.694	1.00	79.65
	MOTA	2332	CB	ALA A 24	9 100.799	31.319	1.964	1.00	80.65
50	MOTA	2333	С	ALA A 24	9 103.232	30.799	2.234	1.00	78.22
	MOTA	2334	0	ALA A 24	9 103.799	30.201	3.150	1.00	78.60
	ATOM	2335	N	ALA A 25	0 103.734	31.878	1.643	1.00	76.27
55	ATOM	2337	CA	ALA A 25	0 104.997	32.476	2.062	1.00	74.32
	ATOM	2338	CB	ALA A 25	0 106.093	32.174	1.042	1.00	75.66
60	MOTA	2339	С	ALA A 25	0 104.840	33.985	2.249	1.00	71.97
	MOTA	2340	0	ALA A 25	0 105.815	34.732	2.160	1.00	72.98
	ATOM	2341	N	ASP A 25	1 103.606	34.422	2.496	1.00	67.46
65	ATOM	2343	CA	ASP A 25	1 103.302	35.836	2.708	1.00	61.94
	MOTA	2344	СВ	ASP A 25	1 101.797	36.090	2.568	1.00	66.00

	MOTA	2345	CG	ASP I	A 2	251	101.277	35.785	1.177	1.00 70.21
5	MOTA	2346	OD1	ASP A	A :	251	101.467	34.647	0.695	1.00 71.96
J	MOTA	2347	OD2	ASP A	A :	251	100.668	36.687	0.566	1.00 72.84
	MOTA	2348	С	ASP A	A :	251	103.760	36.265	4.101	1.00 56.10
10	MOTA	2349	0	ASP 2	A :	251	104.203	37.396	4.298	1.00 56.97
	MOTA	2350	N	ARG A	A :	2 52	103.611	35.364	5.067	1.00 47.53
1.5	MOTA	2352	CA	ARG 2	A :	252	104.009	35.622	6.441	1.00 39.41
15	MOTA	2353	СВ	ARG A	A :	252	102.815	35.472	7.384	1.00 39.71
	ATOM	2354	CG	ARG I	A :	252	101.668	36.426	7.129	1.00 39.66
20	ATOM	2355	CD	ARG .	A :	252	101.975	37.837	7.596	1.00 41.34
	MOTA	2356	NE	ARG	A :	252	100.830	38.714	7.375	1.00 42.20
25	MOTA	2358	CZ	ARG .	A :	252	100.628	39.416	6.263	1.00 44.51
25	ATOM	2359	NH1	ARG .	A :	252	101.498	39.358	5.263	1.00 45.26
	MOTA	2362	NH2	ARG .	A :	252	99.542	40.163	6.141	1.00 44.83
30	ATOM	2365	С	ARG .	A :	252	105.070	34.594	6.798	1.00 34.91
	ATOM	2366	0	ARG .	A :	252	105.315	33.667	6.035	1.00 31.70
35	ATOM	2367	N	MET .	A :	253	105.662	34.730	7.976	1.00 32.83
<i></i>	MOTA	2369	CA	MET	A _.	253	106.697	33.809	8.417	1.00 30.84
-	MOTA	2370	CB	MET	A	253	107.605	34.499	9.441	1.00 30.27
40	MOTA	2371	CG	MET	A	253	108.192	35.807	8.925	1.00 31.26
	MOTA	2372	SD	MET	A	253	109.366	36.637	10.009	1.00 34.11
45	MOTA	2373	CE	MET	A	253	110.335	37.492	8.834	1.00 29.80
43	MOTA	2374	С	MET	A	253	106.130	32.501	8.979	1.00 31.42
	ATOM	2375	0	MET	A	253	106.568	31.419	8.595	1.00 31.42
50	ATOM	2376	N	LAV	A	254	105.144	32.600	9.866	1.00 31.26
	ATOM	2378	CA	VAL	A	254	104.540	31.419	10.480	1.00 32.17
55	ATOM	2379	CB	VAL	A	254	104.781	31.376	12.015	1.00 32.83
	MOTA	2380	CG1	VAL	A	254	106.243	31.143	12.324	1.00 35.48
	ATOM	2381	CG2	VAL	A	254	104.316	32.672	12.658	1.00 33.43
60	ATOM	2382	С	VAL	A	254	103.043	31.378	10.262	1.00 32.00
	ATOM	2383	0	VAL	A	254	102.412	32.410	10.059	1.00 33.43
65	ATOM	2384	N	TYR	A	255	102.478	30.181	10.340	1.00 33.42
<i></i>	ATOM	2386	CA	TYR	A	255	101.044	29.979	10.179	1.00 32.60

	ATOM	2387	СВ	TYR A	A . :	255	100.726	29.448	8.784	1.00	32.98
	ATOM	2388	CG	TYR A	A :	255	100.746	30.482	7.694	1.00	32.36
5	ATOM	2389	CD1	TYR I	A :	255	101.913	30.756	6.987	1.00	31.99
	ATOM	2390	CE1	TYR 2	A	255	101.921	31.674	5.947	1.00	33.10
10	MOTA	2391	CD2	TYR I	A :	255	99.587	31.154	7.335	1.00	31.03
10	MOTA	2392	CE2	TYR 3	A	255	99.583	32.072	6.298	1.00	32.21
	MOTA	2393	CZ	TYR :	A	255	100.750	32.325	5.607	1.00	33.29
15	MOTA	2394	ОН	TYR .	A	255	100.741	33.230	4.573	1.00	38.59
	MOTA	2396	С	TYR .	A	255	100.559	28.955	11.195	1.00	31.73
20	MOTA	2397	0	TYR .	A	255	101.156	27.892	11.330	1.00	32.57
20	МОТА	2398	N	LEU .	A	256	99.490	29.279	11.914	1.00	31.36
	MOTA	2400	CA	LEU .	A	256	98.916	28.370	12.893	1.00	29.13
25	ATOM	2401	CB	LEU	A	256	98.878	28.992	14.280	1.00	28.40
	MOTA	2402	CG	LEU	A	256	100.206	29.125	15.014	1.00	28.89
30	ATOM	2403	CD1	LEU	A	256	101.045	30.208	14.375	1.00	32.20
30	ATOM	2404	CD2	LEU	A	256	99.938	29.462	16.471	1.00	32.12
	ATOM	2405	С	LEU	A	256	97.509	27.984	12.489	1.00	30.86
35	ATOM	2406	0	LEU	A	256	96.748	28.814	11.987	1.00	30.77
	MOTA	2407	N	GLY	A	257	97.203	26.701	12.652	1.00	31.62
40	MOTA	2409	CA	GLY	A	257	95.888	26.180	12.345	1.00	29.36
40	MOTA	2410	С	GLY	A	257	95.287	25.827	13.688	1.00	29.99
	MOTA	2411	0	GLY	A	257	95.662	24.830	14.295	1.00	32.08
45	MOTA	2412	N	LEU	A	258	94.402	26.686	14.177	1.00	29.99
	MOTA	2414	CA	LEU	A	258	93.742	26.507	15.463	1.00	26.58
50	MOTA	2415	СВ	LEU	A	258	93.562	27.871	16.101	1.00	26.90
	ATOM	2416	CG	LEU	A	258	94.866	28.668	16.049	1.00	28.57
	ATOM	2417	CD1	LEU	A	258	94.583	30.144	16.176	1.00	29.14
55	MOTA	2418	CD2	LEU	A	258	95.808	28.177	17.139	1.00	29.78
	MOTA	2419	С	LEU	A	258	92.397	25.838	15.253	1.00	28.27
60	MOTA	2420	0	LEU	A	258	91.404	26.499	14.936	1.00	29.90
50	. ATOM	2421	N	SER	A	259	92.370	24.523	15.434	1.00	28.81
	MOTA	2423	CA	SER	A	259	91.164	23.726	15.229	1.00	30.63
65	MOTA	2424	СВ	SER	A	259	91.491	22.237	15.348	1.00	31.03
	MOTA	2425	OG	SER	A	259	91.783	21.889	16.689	1.00	33.43

	MOTA	2427	С	SER A	259	90.003	24.041	16.152	1.00 31.95
5	MOTA	2428	0	SER A	259	90.184	24.600	17.231	1.00 33.18
3	MOTA	2429	N	ASP A	260	88.808	23.637	15.729	1.00 32.89
	ATOM	2431	CA	ASP A	260	87.616	23.844	16.530	1.00 31.93
10	MOTA	2432	CB	ASP A	260	86.326	23.521	15.739	1.00 33.23
	MOTA	2433	CG	ASP A	260	86.327	22.127	15.080	1.00 32.26
16	MOTA	2434	ODl	ASP A	260	87.138	21.246	15.425	1.00 33.22
15	MOTA	2435	OD2	ASP A	260	85.473	21.904	14.203	1.00 32.09
	MOTA	2436	С	ASP A	260	87.725	23.015	17.807	1.00 30.67
20	ATOM	2437	0	ASP A	260	87.234	23.417	18.856	1.00 34.53
	MOTA	2438	N	TYR A	261	88.414	21.881	17.715	1.00 27.04
25	MOTA	2440	CA	TYR A	261	88.629	20.983	18.847	1.00 26.39
25	MOTA	2441	СВ	TYR A	261	89.490	19.789	18.382	1.00 27.42
	MOTA	2442	CG	TYR A	261	90.064	18.885	19.463	1.00 26.82
30	MOTA	2443	CD1	TYR A	261	91.304	19.157	20.038	1.00 29.26
	MOTA	2444	CE1	TYR A	261	91.855	18.318	21.006	1.00 30.93
35	MOTA	2445	CD2	TYR A	261	89.386	17.745	19.886	1.00 26.82
23	ATOM	2446	CE2	TYR A	261	89.927	16.898	20.853	1.00 28.99
	ATOM	2447	CZ	TYR A	261	91.163	17.194	21.408	1.00 32.15
40	MOTA	2448	OH	TYR A	A 261	91.715	16.368	22.364	1.00 36.84
	MOTA	2450	С	TYR A	A 261	89.345	21.796	19.917	1.00 25.35
45	MOTA	2451	0	TYR A	A 261	88.939	21.824	21.076	1.00 31.74
••	MOTA	2452	N	PHE A	A 262	90.361	22.525	19.488	1.00 23.54
	ATOM	2454	CA	PHE A	A 262	91.151	23.361	20.368	1.00 21.85
50	MOTA	2455	СВ	PHE A	A 262	92.141	24.158	19.514	1.00 19.79
	MOTA	2456	CG	PHE 2	A 262	92.923	25.184	20.275	1.00 20.92
55	MOTA	2457	CD1	PHE I	A 262	93.830	24.806	21.253	1.00 22.00
	MOTA	2458	CD2	PHE	A 262	92.761	26.536	20.002	1.00 20.09
	MOTA	2459	CE1	PHE 2	A 262	94.558	25.759	21.943	1.00 19.37
60	ATOM	2460	CE2	PHE	A 262	93.488	27.497	20.693	1.00 18.46
	MOTA	2461	CZ	PHE 2	A 262	94.386	27.107	21.660	1.00 18.18
65	ATOM	2462	С	PHE 2	A 262	90.244	24.290	21.165	1.00 21.70
UJ.	ATOM	2463	0	PHE :	A 262	90.236	24.266	22.391	1.00 22.47

	MOTA	2464	N	PHE	A	263	89.432	25.060	20.457	1.00	24.02
	ATOM	2466	CA	PHE	A	263	88.522	26.014	21.079	1.00	23.57
5	MOTA	2467	CB	PHE	A	263	87.843	26.845	20.005	1.00	22.46
	MOTA	2468	CG	PHE	A	263	88.783	27.712	19.241	1.00	21.39
10	MOTA	2469	CD1	PHE	A	263	89.462	28.740	19.870	1.00	25.06
10	ATOM	2470	CD2	PHE	A	263	88.982	27.510	17.887	1.00	23.84
	ATOM	2471	CEl	PHE	A	263	90.326	29.555	19.159	1.00	26.31
15	MOTA	2472	CE2	PHE	A	263	89.843	28.319	17.169	1.00	25.34
	MOTA	2473	CZ	PHE	A	263	90.514	29.345	17.804	1.00	25.18
20	MOTA	2474	c .	PHE	A	263	87.457	25.378	21.933	1.00	25.63
20	MOTA	2475	0	PHE	A	263	87.103	25.892	22.992	1.00	29.04
	MOTA	2476	N	ASN	A	264	86.926	24.264	21.463	1.00	27.70
25	ATOM	2478	CA	ASN	A	264	85.882	23.577	22.193	1.00	26.71
	MOTA	2479	CB	ASN	A	264	85.136	22.617	21.278	1.00	26.91
30	MOTA	2480	CG	ASN	A	264	84.183	23.338	20.345	1.00	26.18
20	MOTA	2481	OD1	ASN	A	264	83.115	23.779	20.755	1.00	29.25
	MOTA	2482	ND2	ASN	A	264	84.563	23.462	19.087	1.00	30.59
35	ATOM	2485	С	ASN	A	264	86.359	22.898	23.470	1.00	26.40
	MOTA	2486	0	ASN	A	264	85.587	22.763	24.414	1.00	27.80
40	ATOM	2487	N	THR	A	265	87.628	22.498	23.527	1.00	26.30
	ATOM	2489	CA	THR	A	265	88.154	21.880	24.748	1.00	24.34
	MOTA	2490	CB	THR	A	265	89.572	21.298	24.559	1.00	24.55
45	ATOM	2491	0G1	THR	A	265	90.453	22.296	24.033	1.00	26.59
	MOTA	2493	CG2	THR	A	265	89.532	20.129	23.619	1.00	21.96
50	MOTA	2494	С	THR	A	265	88.175	22.924	25.865	1.00	23.53
	MOTA	2495	0	THR	A	265	88.045	22.602	27.041	1.00	26.65
	MOTA	2496	N	ALA	A	266	88.309	24.186	25.483	1.00	24.00
55	ATOM	2498	CA	ALA	A	266	88.315	25.276	26.437	1.00	22.35
	ATOM	2499	СВ	ALA	A	266	88.582	26.590	25.723	1.00	21.75
60	MOTA	2500	С	ALA	A	266	86.956	25.310	27.113	1.00	23.18
	ATOM	2501	0	ALA	A	266	86.865	25.297	28.336	1.00	.27.51
	ATOM	2502	N	GLY	A	267	85.897	25.286	26.311	1.00	24.69
65	ATOM	2504	CA	GLY	A	267	84.549	25.323	26.852	1.00	24.20
	MOTA	2505	C	GLY	A	267	84.251	24.188	27.812	1.00	27.46

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	MOTA	2506	0	GLY A 267	83.637	24.381	28.869	1.00 27.80
5	MOTA	2507	N	LEU A 268	84.699	22.997	27.445	1.00 28.36
3	MOTA	2509	CA	LEU A 268	84.493	21.816	28.262	1.00 29.18
	MOTA	2510	CB	LEU A 268	84.996	20.580	27.520	1.00 32.75
10	MOTA	2511	CG	LEU A 268	84.728	19.224	28.169	1.00 37.10
	ATOM	2512	CD1	LEU A 268	83.274	18.841	27.978	1.00 38.10
15	ATOM	2513	CD2	LEU A 268	85.633	18.180	27.544	1.00 41.14
15	ATOM	2514	С	LEU A 268	85.209	21.940	29.596	1.00 27.77
	ATOM	2515	0	LEU A 268	84.623	21.696	30.644	1.00 31.78
20	ATOM	2516	N	VAL A 269	86.469	22.344	29.558	1.00 26.94
	MOTA	2518	CA	VAL A 269	87.256	22.478	30.774	1.00 26.89
25	ATOM	2519	СВ	VAL A 269	88.752	22.720	30.448	1.00 24.63
23	MOTA	2520	CG1	VAL A 269	89.491	23.244	31.656	1.00 22.76
	MOTA	2521	CG2	VAL A 269	89.382	21.416	30.007	1.00 21.59
30	MOTA	2522	С	VAL A 269	86.722	23.535	31.740	1.00 28.84
	MOTA	2523	0	VAL A 269	86.687	23.309	32.948	1.00 29.66
35	MOTA	2524	N	TYR A 270	86.298	24.681	31.219	1.00 30.57
33	ATOM	2526	CA	TYR A 270	85.769	25.734	32.080	1.00 28.60
	MOTA	2527	CB	TYR A 270	85.753	27.079	31.350	1.00 28.80
40	ATOM	2528	CG	TYR A 270	87.111	27.734	31.265	1.00 27.47
	MOTA	2529	CD1	TYR A 270	87.766	28.169	32.415	1.00 25.23
45	MOTA	2530	CE1	TYR A 270	89.021	28.754	32.343	1.00 26.36
,,,	MOTA	2531	CD2	TYR A 270	87.750	27.906	30.038	1.00 29.53
	MOTA	2532	CE2	TYR A 270	89.008	28.494	29.959	1.00 25.30
50	MOTA	2533	CZ	TYR A 270	89.632	28.912	31.114	1.00 24.84
	MOTA	2534	ОН	TYR A 270	90.877	29.492	31.047	1.00 30.89
55	MOTA	2536	С	TYR A 270	84.385	25.405	32.631	1.00 29.08
	MOTA	2537	0	TYR A 270	84.034	25.836	33.729	1.00 28.17
	MOTA	2538	N	GLN A 271	83.609	24.628	31.882	1.00 30.32
60	MOTA	2540	CA	GLN A 271	82.269	24.249	32.316	1.00 29.24
	MOTA	2541	СВ	GLN A 271	81.464	23.689	31.139	1.00 27.20
65	MOTA	2542	CG	GLN A 271	79.957	23.680	31.357	1.00 27.60
UJ	MOTA	2543	CD	GLN A 271	79.427	22.387	31.946	1.00 31.03

		MOTA	2544	OE1	GLN	A	271	79.982	21.306	31.724	1.00	35.11
		ATOM	2545	NE2	GLN	A	271	78.318	22.484	32.668	1.00	28.89
	5	ATOM	2548	С	GLN	Α	271	82.329	23.226	33.438	1.00	30.41
		ATOM	2549	0	GLN	A	271	81.839	23.473	34.539	1.00	32.81
	10	MOTA	2550	N	GLU	A	272	82.976	22.098	33.168	1.00	33.61
	10	ATOM	2552	CA	GLU	A	272	83.096	21.011	34.133	1.00	36.97
		MOTA	2553	СВ	GLU	A	272	83.807	19.822	33.492	1.00	40.58
	15	ATOM	2554	CG	GLU	A	272	83.084	19.264	32.274	1.00	48.51
~·		ATOM	2555	CD	GLU	A	272	83.781	18.051	31.674	1.00	55.01
	20	ATOM	2556	OE1	GLU	A	272	84.998	18.129	31.405	1.00	59.64
•	20	MOTA	2557	OE2	GLU	A	272	83.112	17.016	31.468	1.00	57.25
		MOTA	2558	C	GLU	A	272	83.761	21.376	35.460	1.00	36.60
:	25	MOTA	2559	0	GLU	A	272	83.674	20.622	36.425	1.00	40.35
		MOTA	2560	N	ALA	A	273	84.429	22.522	35.512	1.00	35.77
	30	MOTA	2562	CA	ALA	A	273	85.079	22.970	36.741	1.00	34.34
		MOTA	2563	CB	ALA	A	273	86.316	23.798	36.418	1.00	32.87
		MOTA	2564	С	ALA	A	273	84.104	23.778	37.594	1.00	34.54
	35	ATOM	2565	0	ALA	A	273	84.459	24.252	38.679	1.00	35.72
		MOTA	2566	N	GLY	A	274	82.879	23.930	37.086	1.00	35.04
	40	MOTA	2568	CA	GLY	A	274	81.831	24.663	37.779	1.00	35.01
		MOTA	2569	С	GLY	A	274	82.092	26.147	37.930	1.00	34.45
		ATOM	2570	0	GLY	A	274	81.640	26.761	38.897	1.00	36.89
	45	MOTA	2571	N	VAL	A	275	82.752	26.742	36.944	1.00	34.05
		MOTA	2573	CA			275	83.083	28.157	37.018	1.00	34.43
	50	ATOM	2574	CB	VAL	A	275	84.600	28.381	36.778	1.00	35.67
		ATOM	2575	CG1	VAL	A	275	84.898	28.616	35.308	1.00	37.90
		ATOM	2576				275	85.105	29.519	37.638		38.34
	55	MOTA	2577	С			275	82.236	29.078	36.135		
		MOTA	2578	0			275	82.211	30.286	36.343		35.12
	60	MOTA	2579	N			276	81.527	28.523	35.159		32.82
		MOTA	2581	CA			276	80.683	29.352	34.307		29.38
	<i>(</i> -	MOTA	2582	CB			276	80.478	28.695	32.936		27.64
	65	ATOM	2583	CG			276	81.739	28.482	32.082		25.37
		MOTA	2584	CD1	LEU	A	276	81.370	27.883	30.757	1.00	24.34

	MOTA	2585	CD2	LEU A	276	82.459	29.793	31.858	1.00	28.39
•	MOTA	2586	С	LEU A	276	79.363	29.548	35.049	1.00	29.83
5	ATOM	2587	0	LEU A	276	78.323	29.007	34.674	1.00	30.32
	ATOM	2588	N	LYS A	277	79.443	30.271	36.158	1.00	29.10
10	ATOM	2590	CA	LYS A	277	78.294	30.546	37.004	1.00	27.79
	ATOM	2591	СВ	LYS A	277	78.327	29.656	38.246	1.00	29.34
1.5	ATOM	2592	CG	LYS A	277	78.211	28.183	37.956	1.00	33.13
15	ATOM	2593	CD	LYS A	277	78.269	27.369	39.230	1.00	39.28
	MOTA	2594	CE	LYS A	277	78.064	25.888	38.930	1.00	44.23
20	ATOM	2595	NZ	LYS A	277	78.321	25.022	40.119	1.00	46.99
	ATOM	2599	С	LYS A	277	78.386	32.002	37.420	1.00	26.55
25	ATOM	2600	0	LYS A	277	79.474	32.556	37.516	1.00	28.05
23	MOTA	2601	N	MET A	278	77.251	32.608	37.723	1.00	25.90
	ATOM	2603	CA	MET A	278	77.251	33.999	38.111	1.00	26.31
30	ATOM	2604	CB	MET A	278	77.350	34.854	36.864	1.00	25.28
	ATOM	2605	CG	MET A	278	77.496	36.310	37.129	1.00	30.09
35	ATOM	2606	SD	MET A	278	77.760	37.127	35.586	1.00	37.76
55	MOTA	2607	CE	MET A	278	77.452	38.822	36.056	1.00	34.82
	MOTA	2608	С	MET A	278	75.985	34.338	38.872	1.00	28.79
40	MOTA	2609	0	MET A	278	74.910	33.858	38.533	1.00	30.81
	ATOM	2610	N	THR A	279	76.123	35.134	39.925	1.00	30.03
45	MOTA	2612	CA	THR A	279	74.987	35.544	40.728	1.00	29.69
	MOTA	2613	СВ	THR A	279	75.242	35.277	42.211	1.00	29.00
	ATOM	2614	OG1	THR A	279	75.286	33.864	42.431	1.00	31.54
50	ATOM	2616	CG2	THR A	279	74.140	35.863	43.060	1.00	28.67
	MOTA	2617	С	THR A	279	74.714	37.025	40.515	1.00	31.93
55	MOTA	2618	0 .	THR A	279	75.615	37.856	40.644	1.00	32.91
	MOTA	2619	N	LEU A	280	73.482	37.338	40.128	1.00	33.14
	MOTA	2621	CA	LEU A	280	73.062	38.715	39.893	1.00	33.77
60	MOTA	2622	СВ	LEU A	280	72.268	38.826	38.581	1.00	34.16
	MOTA	2623	CG	LEU A	280	72.930	38.757	37.194	1.00	33.72
65	MOTA	2624	CD1	LEU A	280	74.391	38.394	37.271	1.00	33.08
05	MOTA	2625	CD2	LEU A	280	72.186	37.766	36.333	1.00	30.96

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	MOTA	2626	С	LEU	A	280	72.196	39.173	41.064	1.00	35.73 ⁻
	ATOM	2627	0	LEU	Α	280	71.223	38.508	41.438	1.00	33.09
5	MOTA	2628	N	ARG	A	281	72.584	40.290	41.666	1.00	39.95
	ATOM	2630	CA	ARG	Α	281	71.864	40.860	42.796	1.00	42.55
10	ATOM	2631	CB	ARG	Α	281	72.823	41.091	43.975	1.00	44.48
10	ATOM	2632	CG	ARG	A	281	73.472	39.824	44.534	1.00	47.55
	ATOM	2633	CD	ARG	A	281	74.406	40.109	45.712	1.00	50.80
15	MOTA	2634	NE	ARG	A	281	75.660	40.757	45.321	1.00	57.47
	MOTA	2636	CZ	ARG	A	281	76.844	40.145	45.263	1.00	61.45
20	MOTA	2637	NHl	ARG	A	281	76.956	38.852	45.552	1.00	62.83
20	MOTA	2640	NH2	ARG	A	281	77.925	40.825	44.901	1.00	62.95
	MOTA	2643	С	ARG	A	281	71.255	42.190	42.354	1.00	44.72
25	ATOM	2644	0	ARG	A	281	71.782	42.854	41.458	1.00	46.61
	MOTA	2645	N	ASP	A	282	70.179	42.601	43.014	1.00	46.42
30	MOTA	2647	CA	ASP	A	282	69.491	43.849	42.702	1.00	47.87
	ATOM	2648	СВ	ASP	A	282	68.284	44.039	43.634	1.00	49.38
	ATOM	2649	CG	ASP	A	282	67.472	45.280	43.302	1.00	51.57
35	MOTA	2650	OD1	ASP	A	282	66.898	45.349	42.195	1.00	54.37
	MOTA	2651	OD2	ASP	A	282	67.408	46.191	44.150	1.00	54.52
40	MOTA	2652	С	ASP	A	282	70.394	45.080	42.762	1.00	48.67
	ATOM	2653	0	ASP	A	282	70.090	46.089	42.141	1.00	51.13
	ATOM	2654	N	ASP	A	283	71.506	44.996	43.488	1.00	49.71
45	MOTA	2656	CA	ASP	A	283	72.427	46.128	43.608	1.00	50.78
	MOTA	2657	CB	ASP	A	283	73.111	46.138	44.987	1.00	52.25
50	MOTA	2658	CG			283	73.994	44.920	45.224	1.00	55.58
	MOTA	2659		ASP			75.189	44.953	44.847		58.60
	ATOM	2660		ASP			73.498	43.935			58.13
55	ATOM	2661	С			283	73.474	46.236	42.492		51.35
	ATOM	2662	0			283	74.362		42.543		52.74
60	ATOM	2663	N			284	73.395		41.500		51.07
	MOTA	2665	CA			284	74.342		40.399		50.50
	ATOM	2666	СВ			284	74.629			•	48.87
65	ATOM	2667	CG			284	75.169		40.956		46.76
	MOTA	2668	SD	MET	A	284	75.494	41.398	40.353	1.00	43.76

	ATOM	2669	CE	MET A	284	76.804	40.895	41.457	1.00 47.06
5	ATOM	2670	С	MET A	. 284	73.775	46.283	39.289	1.00 53.22
	ATOM	2671	0	MET A	284	74.515	46.824	38.466	1.00 55.85
	ATOM	2672	N	ILE A	285	72.455	46.433	39.285	1.00 55.37
10	MOTA	2674	CA	ILE A	285	71.761	47.252	38.295	1.00 57.52
	ATOM	2675	СВ	ILE A	285	70.287	46.792	38.141	1.00 55.38
16	ATOM	2676	CG2	ILE A	285	69.583	47.611	37.070	1.00 56.70
15	ATOM	2677	CG1	ILE A	285	70.235	45.303	37.783	1.00 53.28
	ATOM	2678	CD1	ILE A	285	68.838	44.723	37.732	1.00 50.41
20	ATOM	2679	С	ILE A	285	71.787	48.707	38.776	1.00 59.50
	ATOM	2680	0	ILE A	285	71.358	48.998	39.889	1.00 60.66
25	ATOM	2681	N	PRO A	286	72.304	49.635	37.951	1.00 61.05
23	ATOM	2682	CD	PRO A	286	72.787	49.459	36.573	1.00 61.08
	MOTA	2683	CA	PRO A	286	72.360	51.048	38.350	1.00 62.59
30	MOTA	2684	CB	PRO A	286	72.862	51.748	37.083	1.00 62.24
	MOTA	2685	CG	PRO P	286	72.481	50.801	35.974	1.00 63.04
35	MOTA	2686	С	PRO A	286	71.004	51.583	38.817	1.00 64.15
55	MOTA	2687	0	PRO A	286	69.986	51.400	38.154	1.00 63.39
	MOTA	2688	N	LYS A	287	71.016	52.268	39.956	1.00 67.25
40	MOTA	2690	CA	LYS A	287	69.813	52.826	40.582	1.00 69.68
	MOTA	2691	CB	LYS A	287	70.204	53.622	41.838	1.00 72.51
45	MOTA	2692	CG	LYS A	287	71.285	54.675	41.607	1.00 75.76
	MOTA	2693	CD	LYS A	A 287	71.483	55.570	42.826	1.00 78.18
	ATOM	2694	CE	LYS A	A 287	72.474	56.697	42.534	1.00 79.24
50	ATOM	2695	NZ	LYS A	A 287	72.603	57.663	43.666	1.00 79.25
	ATOM	2699	С	LYS A	A 287	68.845	53.653	39.724	1.00 69.37
55	ATOM	2700	0	LYS A	A 287	67.735	53.962	40.165	1.00 69.08
	ATOM	2701	N	GLU · I	A 288	69.242	53.996	38.505	1.00 68.49
	ATOM	2703	CA	GLU Z	A 288	68.386	54.792	37.635	1.00 69.17
60	ATOM	2704	CB		A 288	69.152	56.004	37.094	1.00 70.77
	ATOM	2705	CG	GLU Z	A 288	69.897	56.826	38.142	1.00 73.92
65	MOTA	2706	CD	GLU 2	A 288	71.290	56,292	38.461	1.00 76.03
	ATOM	2707	OE1	GLU 2	A 288	71.656	55.195	37.983	1.00 78.84

	MOTA	2708	OE2	GLU	A	288	72.031	56.981	39.196	1.00	76.47
	MOTA	2709	С	GLU	A	288	67.851	53.965	36.469	1.00	68.68
5	MOTA	2710	0	GLU	A	288	68.012	54.345	35.310	1.00	70.05
	ATOM	2711	N	SER	A	289	67.194	52.852	36.772	1.00	68.09
10	MOTA	2713	CA	SER	A	289	66.655	51.985	35.727	1.00	67.82
10	ATOM	2714	CB	SER	A	289	67.516	50.728	35.593	1.00	69.68
	MOTA	2715	OG	SER	A	289	68.868	51.064	35.327	1.00	73.68
15	ATOM	2717	С	SER	A	289	65.202	51.590	35.977	1.00	66.67
	ATOM	2718	0	SER	A	289	64.695	51.730	37.090	1.00	67.98
20	ATOM	2719	N	ALA	A	290	64.537	51.110	34.928	1.00	63.49
20	ATOM	2721	CA	ALA	A	290	63.144	50.691	35.027	1.00	60.33
	ATOM	2722	CB	ALA	A	290	62.525	50.598	33.644	1.00	60.15
25	MOTA	2723	С	ALA	A	290	63.026	49.353	35.744	1.00	58.83
	MOTA	2724	0	ALA	A	290	62.236	49.208	36.673	1.00	59.19
30	ATOM	2725	N	PHE	A	291	63.839	48.390	35.321	1.00	57.41
50	ATOM	2727	CA	PHE	A	291	63.829	47.043	35.894	1.00	55.46
	ATOM	2728	CB	PHE	A	291	64.441	46.044	34.902	1.00	54.51
35	ATOM	2729	CG	PHE	A	291	63.658	45.890	33.622	1.00	52.78
	ATOM	2730	CD1	PHE	A	291	62.493	45.130	33.588	1.00	52.76
40	ATOM	2731	CD2	PHE	A	291	64.085	46.505	32.451	1.00	52.05
	MOTA	2732	CE1	PHE	A	291 '	61.760	44.983	32.414	1.00	51.50
	ATOM	2733	CE2	PHE	A	291	63.360	46.364	31.270	1.00	52.10
45	ATOM	2734	CZ	PHE	A	291	62.194	45.601	31.255	1.00	51.58
	MOTA	2735	C	PHE	A	291	64.519	46.912	37.260	1.00	53.52
50	MOTA	2736	0	PHE	A	291	65.541	47.547	37.524	1.00	53.59
	ATOM	2737	N	ARG	A	292	63.966	46.046	38.104	1.00	50.17
	MOTA	2739	CA	ARG	A	292	64.494	45.799	39.438	1.00	46.66
55	MOTA	2740	CB	ARG	A	292	63.677	46.562	40.481	1.00	46.93
	MOTA	2741	CG	ARG	A	292	63.793	48.065	40.434	1.00	47.97
60	MOTA	2742	CD	ARG	A	292	64.964	48.536	41.256	1.00	49.34
	ATOM	2743	NE	ARG	A	292	65.959	49.215	40.438	1.00	51.19
	MOTA	2745	CZ	ARG	A	292	67.266	48.993	40.521	1.00	52.32
65	MOTA	2746	NH1	ARG	A	292	67.735	48.102	41.385	1.00	52.06
	MOTA	2749	NH2	ARG	A	292	68.106	49.674	39.750	1.00	52.32

	ATOM	2752	С	ARG A	A :	292	64.317	44.322	39.715	1.00	46.15
5	ATOM	2753	0	ARG A	A :	292	63.333	43.720	39.278	1.00	47.65
3	ATOM	2754	N	LEU Z	A :	293	65.278	43.725	40.409	1.00	44.69
	ATOM	2756	CA	LEU Z	A :	293	65.169	42.320	40.769	1.00	42.76
10	ATOM	2757	CB	LEU 2	A :	293	66.546	41.648	40.819	1.00	37.95
	MOTA	2758	CG	LEU 2	A :	293	67.252	41.569	39.468	1.00	34.56
15	MOTA	2759	CD1	LEU 2	A :	293	68.649	41.034	39.630	1.00	33.82
15	MOTA	2760	CD2	LEU .	A :	293	66.459	40.692	38.537	1.00	32.43
	MOTA	2761	С	LEU .	ΑÌ	293	64.516	42.349	42.141	1.00	44.58
20	ATOM	2762	0	LEU .	A :	293	65.169	42.154	43.164	1.00	46.89
	ATOM	2763	N	THR .	Α :	294	63.239	42.716	42.150	1.00	44.93
25	MOTA	2765	CA	THR .	A.	294	62.453	42.804	43.370	1.00	45.20
25	ATOM	2766	CB	THR	A	294	62.220	44.271	43.781	1.00	44.99
	MOTA	2767	OG1	THR	A	294	61.502	44.955	42.748	1.00	45.39
30	ATOM	2769	CG2	THR	A	294	63.547	44.983	44.012	1.00	46.79
	ATOM	2770	С	THR	A	294	61.111	42.175	43.055	1.00	46.49
35	ATOM	2771	0	THR	A	294	60.639	42.269	41.922	1.00	48.98
33	ATOM	2772	N	THR	A	295	60.500	41.520	44.035	1.00	47.36
	ATOM	2774	CA	THR	A	295	59.204	40.892	43.812	1.00	48.40
40	ATOM	2775	CB	THR	A	295	58.720	40.139	45.063	1.00	47.45
	ATOM	2776	OG1	THR	A	295	58.837	40.986	46.210	1.00	47.38
45	ATOM	2778	CG2	THR	A	295	59.550	38.891	45.287	1.00	47.66
45	MOTA	2779	С	THR	A	295	58.179	41.952	43.410	1.00	49.59
	MOTA	2780	0	THR	A	295	57.292	41.693	42.599	1.00	50.18
50	MOTA	2781	N	SER	A	296	58.347	43.157	43.950	1.00	51.29
	MOTA	2783	CA	SER	A	296	57.468	44.288	43.667	1.00	52.78
55	MOTA	2784	СВ	SER	A	296	57.978	45.538	44.390	1.00	54.54
55	MOTA	2785	QG	SER	A	296	57.219	46.688	44.056	1.00	55.48
	ATOM	2787	С	SER	A	296	57.412	44.551	42.172	1.00	52.76
60	ATOM	2788	0	SER	A	296	56.340	44.532	41.562	1.00	52.27
	ATOM	2789	N	PHE	A	297	58.579	44.779	41.583	1.00	52.83
65	ATOM	2791	CA	PHE	A	297	58.656	45.033	40.160	1.00	53.94
O)	ATOM	2792	СВ	PHE	A	297	60.093	45.286	39.730	1.00	55.28

						154				
	MOTA	2793	CG	PHE A	297	60.217	45.651	38.293	1.00	57.93 ⁻
	MOTA	2794	CD1	PHE A	297	60.028	46.964	37.884	1.00	57.66
5	ATOM	2795	CD2	PHE A	297	60.459	44.673	37.334	1.00	59.49
	MOTA	2796	CE1	PHE A	297	60.072	47.299	36.539	1.00	59.28
10	MOTA	2797	CE2	PHE A	297	60.503	44.997	35.990	1.00	59.71
10	MOTA	2798	CZ	PHE A	297	60.309	46.315	35.590	1.00	59.93
	MOTA	2799	С	PHE A	297	58.090	43.837	39.408	1.00	54.31
15	ATOM	2800	0	PHE A	297	57.245	43.990	38.530	1.00	54.87
	MOTA	2801	N	PHE A	298	58.542	42.643	39.776	1.00	55.31
20	MOTA	2803	CA	PHE A	298	58.058	41.420	39.149	1.00	55.92
20	ATOM	2804	СВ	PHE A	298	58.793	40.199	39.709	1.00	52.95
	ATOM	2805	CG	PHE A	298	60.118	39.943	39.057	1.00	49.79
25	MOTA	2806	CD1	PHE A	298	61.094	40.934	39.019	1.00	46.14
	ATOM	2807	CD2	PHE A	298	60.384	38.716	38.463	1.00	47.97
30	ATOM	2808	CEl	PHE A	298	62.312	40.709	38.400	1.00	.44.36
50	ATOM	2809	CE2	PHE A	298	61.600	38.482	37.842	1.00	46.94
	ATOM	2810	CZ	PHE A	298	62.567	39.482	37.810	1.00	45.78
35	ATOM	2811	С	PHE A	298	56.556	41.274	39.364	1.00	58.12
	ATOM	2812	0	PHE A	298	55.904	40.453	38.716	1.00	58.39
40	ATOM	2813	N	GLY A	299	56.019	42.067	40.287	1.00	60.34
	MOTA	2815	CA	GLY A	299	54.599	42.042	40.576	1.00	62.61
	ATOM	2816	С	GLY A	299	53.777	42.541	39.404	1.00	64.03
45	MOTA	2817	0	GLY A	299	52.622	42.156	39.236	1.00	64.07
	MOTA	2818	N	THR A	300	54.377	43.379	38.571	1.00	65.39
50	MOTA	2820	CA	THR A	300	53.672	43.904	37.416	1.00	67.18
	MOTA	2821	CB	THR A	300	54.532	44.935	36.658	1.00	66.57
	MOTA	2822	OG1	THR A	300	55.806	44.361	36.349	1.00	67.61
55	MOTA	2824	CG2	THR A	300	54.736	46.181	37.503	1.00	67.44
	MOTA	2825	С	THR A	300	53.258	42.785	36.460	1.00	69.21
60	MOTA	2826	0	THR A	300	52.234	42.896	35.777	1.00	72.16
	MOTA	2827	N	PHE A	301	54.055	41.774	36.384	1.00	69.34
	ATOM	2829	CA	PHE A	301	53.704	40.675	35.483	1.00	69.26
65	MOTA	2830	СВ	PHE A	301	54.608	40.680	34:241	1.00	66.47
	MOTA	2831	CG	PHE A	301	56.070	40.882	34.532	1.00	64.06

MOTA

2867

CA

VAL A 305

55.955 36.809

43.226

1.00 74.85

PCT/US98/13007 WO 98/58961

CD1 PHE A 301 MOTA 2832 56.629 42.154 CD2 PHE A 301 39.806 34.838 MOTA 2833 56.895 5 CE1 PHE A 301 34.716 MOTA 2834 57.983 42.354 ATOM 2835 CE2 PHE A 301 58.251 39.995 35.077 10 MOTA 2836 ÇZ PHE A 301 58.795 MOTA 2837 C PHE A 301 53.635 PHE A 301 53.032 MOTA 2838 0 15 MOTA 2839 N LEU A 302 54.199 39.072 37.250 MOTA 2841 CA LEU A 302 54.198 37.748 37.866 20 MOTA 2842 CB LEU A 302 55.573 37.098 37.698 ATOM 2843 CG **LEU A 302** 55.833 36.472 36.337 MOTA 2844 CD1 LEU A 302 57.217 35.860 25 CD2 LEU A 302 54.767 35.425 36.070 MOTA 2845 MOTA 2846 С LEU A 302 53.782 37.667 39.338 30 **ATOM** 2847 0 LEU A 302 54.567 37.228 40.173 52.551 **ATOM** 2848 N PRO A 303 38.075 39.678 ATOM PRO A 303 51.687 39.012 38.927 2849 CD 35 ATOM 2850 CA PRO A 303 52.127 38.002 41.087 MOTA 2851 PRO A 303 51.725 39.440 41.359 CB 40 MOTA 2852 CG PRO A 303 50.963 39.803 40.046 PRO A 303 MOTA 2853 С 50.924 37.070 41.346 **ATOM** 2854 PRO A 303 49.809 37.403 40.953 0 45 **ATOM** 2855 N **GLU A 304** 51.087 35.893 41.943 **ATOM** 2857 CA **GLU A 304** 52.308 35.254 42.438 50 MOTA 2858 CB **GLU A 304** 52.767 34.203 41.437 MOTA 2859 CG **GLU A 304** 51.698 33.132 41.207 32.502 42.499 ATOM 2860 CD **GLU A 304** 51.179 55 MOTA 2861 OE1 GLU A 304 32.923 42.977 50.103

135 34.478 1.00 62.60 1.00 63.14 1.00 61.30 1.00 61.28 41.274 35.016 1.00 61.18 39.268 36.062 1.00 71.49 38.380 35.458 1.00 71.98 1.00 74.67 1.00 77.56 1.00 75.96 1.00 75.33 36.332 1.00 75.78 1.00 74.63 1.00 80.50 1.00 81.63 1.00 82.47 1.00 83.87 1.00 82.85 1.00 83.29 1.00 84.24 1.00 82.81 1.00 84.47 1.00 81.68 1.00 80.61 1.00 83.61 1.00 86.52 1.00 88.36 1.00 89.84 1.00 89.41 MOTA 2862 OE2 GLU A 304 51.847 31.594 43.036 60 MOTA 2863 C **GLU A 304** 53.501 35.887 43.175 1.00 78.52 MOTA 2864 0 GLU A 304 53.383 36.142 44.361 1.00 78.12 MOTA 2865 N VAL A 305 54.817 36.106 42.616 1.00 76.19 65

	MOTA	2868	CB	VAL	A	305	56.904	37.447	42.170	1.00	74.41
ŧ	MOTA	2869	CG1	VAL	A	305	57.405	36.392	41.202	1.00	74.39
5	MOTA	2870	CG2	VAL	A	305	56.218	38.578	41.433	1.00	75.20
	MOTA	2871	С	VAL	A	305	55.622	37.833	44.316	1.00	74.68
10	MOTA	2872	0	VAL	A	305	56.110	37.711	45.438	1.00	75.57
10	MOTA	2873	N	ALA	A	306	54.746	38.790	44.011	1.00	73.53
	MOTA	2875	CA	ALA	A	306	54.357	39.823	44.975	1.00	71.77
15	ATOM	2876	CB	ALA	Α	306	53.893	41.068	44.248	1,00	71.20
	ATOM	2877	С	ALA	A	306	53.273	39.348	45.935	1.00	71.50
	ATOM	2878	0	ALA	A	306	53.087	39.924	47.005	1.00	70.71
20	ATOM	2879	N	LYS	A	307	52.563	38.300	45.529	1.00	71.78
•	ATOM	2881	CA	LYS	A	307	51.474	37.697	46.300	1.00	71.82
25	ATOM	2882	CB	LYS	A	307	50.508	37.010	45.324	1.00	72.68
	MOTA	2883	CG	LYS	A	307	49.332	36.270	45.944	1.00	75.23
20	MOTA	2884	CD	LYS	A	307	48.473	35.638	44.848	1.00	76.97
30	MOTA	2885	CE	LYS	A	307	47.247	34.935	45.413	1.00	77.90
	MOTA	2886	NZ	LYS	A	307	46.352	34.434	44.331	1.00	77.64
35	MOTA	2890	С	LYS	A	307	51.946	36.698	47.374	1.00	71.04
	MOTA	2891	0	LYS	A	307	51.393	36.651	48.473	1.00	72.05
40	MOTA	2892	N	LYS	A	308	52.958	35.898	47.050	1.00	69.00
40	MOTA	2894	CA	LYS	A	308	53.480	34.903	47.979	1.00	66.59
	MOTA	2895	СВ	LYS	A	308	53.891	33.636	47.219	1.00	70.09
45	ATOM	2896	CG	LYS	A	308	53.887	32.346	48.046	1.00	75.24
	ATOM	2897	CD	LYS	A	308	54.920	32.351	49.172	1.00	79.31
50	MOTA	2898	CE	LYS	A	308	54.842	31.091	50.027	1.00	81.43
	ATOM	2899	NZ	LYS	A	308	55.760	31.141	51.206	1.00	82.05
	ATOM	2903	С	LYS	A	308	54.680	35.467	48.722	1.00	64.17
55	MOTA	2904	0	LYS	A	308	54.840	35.240	49.921	1.00	65.45
	ATOM	2905	N	PHE	A	309	55.526	36.200	48.007	1.00	60.99
60	MOTA	2907	CA	PHE	A	309	56.725	36.784	48.597	1.00	57.85
00	MOTA	2908	CB	PHE	A	309	57.974	36.274	47.853	1.00	53.07
	MOTA	2909	CG	PHE	A	309	58.085	34.762	47.808	1.00	45.45
65	MOTA	2910	CD1	PHE	A	309	58.551	34.050	48.906	1.00	42.52
	ATOM	2911	CD2	PHE	A	309	57.689	34.052	46.679	1.00	43.68

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	MOTA	2912	CE1	PHE A 309	58.619	32.655	48.883	1.00 38.70
5	ATOM	2913	CE2	PHE A 309	57.754	32.655	46.647	1.00 42.05
3	MOTA	2914	CZ	PHE A 309	58.219	31.959	47.753	1.00 39.93
	ATOM	2915	С	PHE A 309	56.616	38.312	48.553	1.00 58.92
10	ATOM	2916	0	PHE A 309	57.266	38.978	47.743	1.00 58.89
	MOTA	2917	N	PRO A 310	55.822	38.885	49.472	1.00 60.26
1.5	MOTA	2918	CD	PRO A 310	55.173	38.142	50.569	1.00 61.34
15	MOTA	2919	CA	PRO A 310	55.560	40.320	49.613	1.00 61.10
	MOTA	2920	СВ	PRO A 310	54.522	40.358	50.736	1.00 62.64
20	MOTA	2921	CG	PRO A 310	54.934	39.217	51.598	1.00 61.52
	ATOM	2922	С	PRO A 310	56.737	41.239	49.927	1.00 61.05
25	ATOM	2923	0	PRO A 310	57.307	41.187	51.016	1.00 61.70
23	ATOM	2924	N	ASN A 311	57.044	42.121	48.979	1.00 62.21
	ATOM	2926	CA	ASN A 311	58.116	43.111	49.111	1.00 63.11
30	ATOM	2927	CB	ASN A 311	57.667	44.247	50.037	1.00 66.59
	ATOM	2928	CG	ASN A 311	58.589	45.448	49.976	1.00 69.63
35	ATOM	2929	OD1	ASN A 311	58.596	46.186	48.991	1.00 71.72
55	ATOM	2930	ND2	ASN A 311	59.373	45.651	51.027	1.00 71.51
	MOTA	2933	C	ASN A 311	59.458	42.547	49.583	1.00 61.81
40	MOTA	2934	0	ASN A 311	59.855	42.737	50.738	1.00 63.25
	MOTA	2935	N	MET A 312	60.155	41.870	48.673	1.00 58.63
45	MOTA	2937	CA	MET A 312	61.459	41.266	48.950	1.00 53.64
	ATOM	2938	СВ	MET A 312	61.316	39.768	49.269	1.00 53.53
	ATOM	2939	CG	MET A 312	60.599	39.405	50.570	1.00 52.28
50	ATOM	2940	SD	MET A 312	60.259	37.617	50.633	1.00 52.18
	ATOM	2941	CE	MET A 312	58.896	37.534	51.788	1.00 54.70
55	ATOM	2942	С	MET A 312	62.320	41.411	47.699	1.00 49.72
	MOTA	2943	0	MET A 312	61.801	41.656	46.605	1.00 49.26
	MOTA	2944	N	LYS A 313	63.630	41.275	47.865	1.00 44.80
60	ATOM	2946	CA	LYS A 313	64.549	41.363	46.741	1.00 41.81
	MOTA	2947	CB	LYS A 313	65.952	41.707	47.230	1.00 42.38
65	MOTA	2948	CG	LYS A 313	66.050	43.081	47.846	1.00 47.04
33	MOTA	2949	CD	LYS A 313	65.762	44.159	46.818	1.00 49.35

	ATOM	2950	CE	LYS A 31	.3	65.599	45.515	47.481	1.00	53.10-
	ATOM	2951	NZ	LYS A 31	.3	66.805	45.897	48.268	1.00	56.75
5	ATOM	2955	С	LYS A 31	.3	64.562	40.018	46.026	1.00	39.61
	MOTA	2956	0	LYS A 31	.3	64.133	39.013	46.592	1.00	38.93
10	MOTA	2957	N	ILE A 31	L 4	65.040	40.005	44.786	1.00	37.30
10	MOTA	2959	CA	ILE A 31	.4	65.115	38.785	43.995	1.00	34.17
	MOTA	2960	СВ	ILE A 31	.4	64.224	38.865	42.716	1.00	33.93
15	MOTA	2961	CG2	ILE A 31	.4	64.607	37.783	41.708	1.00	32.86
	MOTA	2962	CG1	ILE A 31	.4	62.745	38.728	43.093	1.00	33.03
20	MOTA	2963	CD1	ILE A 31	14	61.824	38.459	41.920	1.00	32.90
20	MOTA	2964	С	ILE A 31	4	66.567	38.565	43.604	1.00	33.46
	MOTA	2965	0	ILE A 31	14	67.272	39.512	43.271	1.00	32.77
25	MOTA	2966	N	GLN A 31	15	67.024	37.321	43.698	1.00	32.44
	MOTA	2968	CA	GLN A 31	15	68.391	36.975	43.336	1.00	32.66
30	MOTA	2969	СВ	GLN A 31	L 5	69.118	36.366	44.529	1.00	34.35
50	MOTA	2970	CG	GLN A 31	15	70.624	36.305	44.376	1.00	35.08
	MOTA	2971	CD	GLN A 31	15	71.317	35.839	45.639	1.00	33.95
35	MOTA	2972	OEI	GLN A 31	L 5	71.579	36.620	46.548	1.00	34.44
	MOTA	2973	NE2	GLN A 31	15	71.607	34.554	45.701	1.00	37.53
40	ATOM	2976	С	GLN A 31	15	68.307	35.964	42.207	1.00	33.04
	MOTA	2977	0	GLN A 31	15	67.520	35.024	42.271	1.00	35.05
	ATOM	2978	N	ILE A 31	16	69.116	36.155	41.175	1.00	34.76
45	MOTA	2980	CA	ILE A 31	16	69.106	35.268	40.018	1.00	33.81
	ATOM	2981	CB	ILE A 31	16	68.711	36.067	38.750	1.00	35.72
50	ATOM	2982	CG2	ILE A 31	16	68.944	35.247	37.498	1.00	37.16
	MOTA	2983	CG1	ILE A 31	16	67.249	36.518	38.864	1.00	36.98
	MOTA	2984	CD1	ILE A 3	16	66.695	37.179	37.633	1.00	38.34
55	MOTA	2985	С	ILE A 3	16	70.445	34.548	39.823	1.00	32.92
	MOTA	2986	0	ILE A 3	16	71.513	35.160	39.900	1.00	33.03
60	MOTA	2987	N .	HIS A 3	17	70.386	33.238	39.609	1.00	33.55
	MOTA	2989	CA	HIS A 3	17	71.586	32.428	39.409	1.00	32.00
	MOTA	2990	СВ	HIS A 3	17	71.532	31.162	40.264	1.00	32.50
65	MOTA	2991	CG	HIS A 3	17	71.659	31.411	41.735	1.00	32.14
	MOTA	2992	CD2	HIS A 3	17	71.889	32.548	42.435	1.00	31.63

	MOTA	2993	ND1	HIS	A	317	71.572	30.398	42.666	1.00	32.81
5	ATOM	2995	CE1	HIS	A	317	71.746	30.899	43.875	1.00	33.94
3	MOTA	2996	NE2	HIS	A	317	71.941	32.201	43.761	1.00	32.58
	ATOM	2998	С	HIS	A	317	71.720	32.014	37.953	1.00	33.04
10	MOTA	2999	0	HIS	A	317	70.844	31.335	37.416	1.00	33.39
	ATOM	3000	N	VAL	A	318	72.842	32.377	37.344	1.00	33.42
15	MOTA	3002	CA	VAL	A	318	73.129	32.059	35.952	1.00	31.61
15	ATOM	3003	СВ	VAL	A	318	73.535	33.338	35.193	1.00	29.43
	MOTA	3004	CG1	VAL	A	318	73.993	33.003	33.789	1.00	30.90
20	ATOM	3005	CG2	VAL	A	318	72.376	34.311	35.161	1.00	27.51
	MOTA	3006	С	VAL	A	318	74.273	31.043	35.861	1.00	32.81
25	ATOM	3007	0	VAL	A	318	75.392	31.325	36.296	1.00	33.30
23	ATOM	3008	N	SER	A	319	73.990	29.858	35.329	1.00	32.14
	ATOM	3010	CA	SER	A	319	75.021	28.836	35.175	1.00	32.12
30	MOTA	3011	CB	SER	A	319	74.919	27.778	36.271	1.00	28.41
	MOTA	3012	OG	SER	A	319	73.723	27.039	36.158	1.00	29.90
35	MOTA	3014	С	SER	A	319	74.900	28.170	33.815	1.00	33.52
33	MOTA	3015	0	SER	A	319	73.834	28.174	33.210	1.00	37.50
	MOTA	3016	N	ALA	A	320	76.005	27.632	33.319	1.00	35.33
40	MOTA	3018	CA	ALA	A	320	76.009	26.958	32.031	1.00	35.02
	MOTA	3019	CB	ALA	A	320	77.417	26.914	31.461	1.00	36.41
45	MOTA	3020	С	ALA	A	320	75.466	25.547	32.210	1.00	36.23
.5	MOTA	3021	0	ALA	A	320	75.955	24.786	33.047	1.00	35.14
	ATOM	3022	N	SER	A	321	74.459	25.204	31.415	1.00	38.15
50	ATOM	3024	CA	SER	A	321	73.835	23.891	31.478	1.00	38.99
	ATOM	3025	CB	SER	A	321	72.472	23.935	30.786	1.00	38.78
55	ATOM	3026	OG	SER	A	321	72.418	24.984	29.834	1.00	40.20
	ATOM	3028	С	SER	A	321	74.724	22.824	30.848	1.00	40.78
	ATOM	3029	0	SER	A	321	74.942	21.760	31.430	1.00	44.24
60	ATOM	3030	N	THR	A	322	75.249	23.126	29.666	1.00	39.66
	MOTA	3032	CA	THR	A	322	76.119	22.214	28.939	1.00	36.30
65	ATOM	3033	CB	THR	A	322	75.355	21.553	27.750	1.00	39.80
5 5	MOTA	3034	OG1	THR	A	322	74.613	22.548	27.027	1.00	40.24

	ATOM	3036	CG2	THR	A	322	74.398	20.473	28.253	1.00	41.99
	ATOM	3037	С	THR	A	322	77.332	22.985	28.416		33.41
5	ATOM	3038	0	THR	A	322	77.372	24.214	28.487		31.80
	ATOM	3039	N	PRO			78.378	22.272	27.966		32.97
	ATOM	3040	CD	PRO	A	323	78.606	20.826	28.133		31.42
10	ATOM	3041	CA	PRO	A	323	79.581	22.925	27.439	1.00	31.79
	ATOM	3042	CB	PRO	A	323	80.471	21.739	27.089	1.00	31.21
15	ATOM	3043	CG	PRO	A	323	80.099	20.740	28.128	1.00	31.27
	ATOM	3044	С	PRO	A	323	79.279	23.757	26.191	1.00	33.72
•	ATOM	3045	0	PRO	Ą	323	78.727	23.237	25.218	1.00	35.68
20	ATOM	3046	N	PRO	A	324	79.650	25.053	26.196	1.00	34.41
	ATOM	3047	CD	PRO	A	324	80.303	25.799	27.286	1.00	34.36
25	ATOM	3048	CA	PRO	A	324	79.407	25.933	25.050	1.00	36.18
	MOTA	3049	СВ	PRO	A	324	79.863	27.301	25.567	1.00	34.56
30	MOTA	3050	CG	PRO	A	324	80.913	26.963	26.551	1.00	34.54
30	ATOM	3051	С	PRO	A	324	80.186	25.511	23.811	1.00	39.38
	ATOM	3052	0	PRO	A	324	81.397	25.278	23.871	1.00	43.04
35	ATOM	3053	N	HIS	A	325	79.468	25.364	22.704	100	39.27
	ATOM	3055	CA	HIS	A	325	80.064	24.973	21.434	1.00	39.40
40	ATOM	3056	CB	HIS	A	325	79.045	24.236	20.562	1.00	44.72
	MOTA	3057	CG	HIS	A	325	78.817	22.813	20.955	1.00	49.32
	MOTA	3058	CD2	HIS	A	325	79.471	22.015	21.831	1.00	51.91
45	ATOM	3059	ND1	HIS	A	325	77.817	22.041	20.405	1.00	52.21
	MOTA	3061	CE1	HIS	A	325	77.864	20.827	20.924	1.00	53.83
50	MOTA	3062	NE2	HIS	A	325	78.859	20.785	21.793	1.00	54.25
	MOTA	3064	С	HIS	A	325	80.537	26.183	20.658	1.00	37.89
	MOTA	3065	0	HIS	A	325	80.011	27.285	20.825	1.00	39.24
55	MOTA	3066	N	LEU	A	326	81.514	25.958	19.788	1.00	36.12
	ATOM	3068	CA	LEU	A	326	82.056	26.993	18.928	1.00	34.45
60	MOTA	3069	СВ	LEU	A	326	83.370	27.535	19.487	1.00	34.16
	MOTA	3070	CG	LEU	A	326	83.906	28.758	18.745	1.00	34.49
	MOTA	3071	CD1	LEU	A	326	82.912	29.878	18.854	1.00	36.18
65	ATOM	3072	CD2	LEU	A	326	85.220	29.194	19.321	1.00	38.20
	MOTA	3073	С	LEU	A	326	82.285	26.329	17.569	1.00	34.87

	MOTA	3074	0	LEU A 326	82.981	25.311	17.469	1.00 3	5.31
	ATOM	3075	N	SER A 327	81.639	26.863	16.542	1.00 3	2.90
5	ATOM	3077	CA	SER A 327	81.755	26.323	15.202	1.00 3	2.76
	MOTA	3078	СВ	SER A 327	80.359	26.173	14.591	1.00 3	4.44
10	ATOM	3079	OG	SER A 327	80.390	25.407	13.394	1.00 4	1.05
	ATOM	3081	С	SER A 327	82.607	27.247	14.343	1.00 3	0.57
	ATOM	3082	0	SER A 327	82.391	28.458	14.329	1.00 3	1.41
15	ATOM	3083	N	VAL A 328	83.601	26.691	13.658	1.00 2	9.84
	ATOM	3085	CA	VAL A 328	84.452	27.502	12.795	1.00 2	8.60
20	ATOM	3086	СВ	VAL A 328	85.951	27.275	13.056	1.00 2	5.61
	ATOM	3087	CG1	VAL A 328	86.733	28.389	12.423	1.00 2	4.98
	ATOM	3088	CG2	VAL A 328	86.245	27.218	14.544	1.00 2	3.23
25	ATOM	3089	С	VAL A 328	84.159	27.174	11.336	1.00 3	0.19
	ATOM	3090	0	VAL A 328	84.221	26.015	10.928	1.00 2	9.56
30	ATOM	3091	N	GLN A 329	83.791	28.194	10.569	1.00 3	2.32
	ATOM	3093	CA	GLN A 329	83.479	28.038	9.150	1.00 3	4.28
35	ATOM	3094	СВ	GLN A 329	81.956	27.972	8.939	1.00 3	9.33
33	MOTA	3095	CG	GLN A 329	81.222	26.924	9.774	1.00 4	8.55
	MOTA	3096	CD	GLN A 329	80.246	27.532	10.785	1.00 5	3.31
40	ATOM	3097	OE1	GLN A 329	79.321	26.860	11.245	1.00 5	7.70
	ATOM	3098	NE2	GLN A 329	80.456	28.796	11.142	1.00 5	3.89
45	ATOM	3101	С	GLN A 329	84.050	29.259	8.416	1.00 3	12.74
45	ATOM	3102	0	GLN A 329	84.337	30.279	9.048	1.00 3	34.32
	ATOM	3103	N	PRO A 330	84.176	29.193	7.073	1.00 3	30.85
50	ATOM	3104	CD	PRO A 330	83.896	28.019	6.226	1.00 2	27.63
	ATOM	3105	CA	PRO A 330	84.711	30.301	6.265	1.00 3	30.72
55	ATOM	3106	CB	PRO A 330	84.597	29.765	4.842	1.00 2	27.09
50	ATOM	3107	CG	PRO A 330	84.754	28.286	5.026	1.00 2	27.10
	ATOM	3108	С	PRO A 330	83.968	31.635	6.416	1.00	33.05
60	MOTA	3109	0	PRO A 330	84.501	32.695	6.084	1.00	34.38
	ATOM	3110	N	THR A 331	82.733	31.575	6.904	1.00	36.91
65	MOTA	3112	CA	THR A 331	81.918	32.767	7.106	1.00	39.55
05	ATOM	3113	СВ	THR A 331	80.414	32.414	7.065	1.00	41.23

		•								
	MOTA	3114	OG1	THR A	331	80.179	31.221	7.826	1.00	42.58-
	ATOM	3116	CG2	THR A	331	79.959	32.180	5.630	1.00	42.18
5	ATOM	3117	C .	THR A	331	82.244	33.469	8.425	1.00	41.21
	MOTA	3118	0	THR A	331	82.114	34.691	8.535	1.00	44.22
10	ATOM	3119	N	GLY A	332	82.671	32.691	9.418	1.00	39.90
10	ATOM	3121	CA	GLY A	332	83.005	33.240	10.720	1.00	36.14
	MOTA	3122	С	GLY A	332	82.788	32.188	11.788	1.00	36.29
15	MOTA	3123	0	GLY A	332	82.399	31.058	11.472	1.00	36.21
	ATOM	3124	N	LEU A	333	83.048	32.542	13.044	1.00	35.70
20	MOTA	3126	CA	LEU A	333	82.875	31.606	14.153	1.00	34.12
20	MOTA	3127	CB	LEU A	333	84.037	31.704	15.157	1.00	31.15
	MOTA	3128	CG	LEU A	333	85.438	32.183	14.766	1.00	30.79
25	ATOM	3129	CD1	LEU A	333	86.402	31.841	15.886	1.00	29.93
	MOTA	3130	CD2	LEU A	333	85.908	31.539	13.492	1.00	32.11
30	ATOM	3131	С	LEU A	333	81.570	31.889	14.889	1.00	34.02
<i>3</i> 0	MOTA	3132	0	LEU A	333	81.283	33.037	15.233	1.00	35.28
	MOTA	3133	N	THR A	334	80.777	30.850	15.123	1.00	32.89
35	MOTA	3135	CA	THR A	334	79.521	30.997	15.842	1.00	30.97
	ATOM	3136	CB	THR A	334	78.344	30.435	15.037	1.00	30.40
40	MOTA	3137	OG1	THR A	334	78.673	29.124	14.568	1.00	36.09
,,	ATOM	3139	CG2	THR A	334	78.043	31.324	13.850	1.00	27.87
	ATOM	3140	С	THR A	334	79.649	30.274	17.184	1.00	30.59
45	ATOM	3141	0	THR A	334	80.293	29.229	17.267	1.00	31.11
	ATOM	3142	N	PHE A	335	79.052	30.853	18.225	1.00	31.55
50	ATOM	3144	CA	PHE A	335	79.095	30.334	19.598	1.00	30.95
	MOTA	3145	CB _.	PHE A	335	79.579	31.465	20.530	1.00	31.73
	ATOM	3146	CG	PHE A	335	80.138	31.001	21.853	1.00	32.43
55	ATOM	3147	CD1	PHE A	335	81.224	30.134	21.907	1.00	34.10
	MOTA	3148	CD2	PHE A	335	79.611	31.483	23.044	1.00	33.32
60	ATOM	3149	CE1	PHE A	335	81.779	29.755	23.124	1.00	36.27
	ATOM	3150	CE2	PHE A	335	80.156	31.113	24.267	1.00	36.93
	ATOM	3151	cz	PHE A	335	81.243	30.248	24.308	1.00	36.83
65	MOTA	3152	С	PHE A	335	77.674	29.910	19.975	1.00	29.88
	MOTA	3153	0	PHE A	335	76.722	30.616	19.666	1.00	29.27

	ATOM	3154	N	TYR A 33	6 77.533	28.779	20.658	1.00	30.87
5	MOTA	3156	CA	TYR A 33	6 76.216	28.281	21.052	1.00	33.87
3	ATOM	3157	CB	TYR A 33	6 75.936	26.953	20.345	1.00	35.86
	MOTA	3158	CG	TYR A 33	6 76.081	27.036	18.844	1.00	36.58
10	ATOM	3159	CD1	TYR A 33	6 75.107	27.665	18.072	1.00	38.45
	ATOM	3160	CE1	TYR A 33	6 75.245	27.791	16.700	1.00	38.32
15	ATOM	3161	CD2	TYR A 33	6 77.207	26.526	18.198	1.00	34.97
15	ATOM	3162	CE2	TYR A 33	6 77.357	26.647	16.819	1.00	36.42
	ATOM	3163	CZ	TYR A 33	6 76.369	27.285	16.078	1.00	37.51
20	ATOM	3164	ОН	TYR A 33	6 76.494	27.435	14.715	1.00	41.83
	ATOM	3166	С	TYR A 33	6 76.104	28.096	22.565	1.00	36.66
25	ATOM	3167	0	TYR A 33	6 76.280	26.988	23.074	1.00	39.11
25	ATOM	3168	N	PRO A 33	7 75.858	29.187	23.309	1.00	36.71
	ATOM	3169	CD	PRO A 33	7 75.890	30.605	22.904	1.00	36.11
30	MOTA	3170	CA	PRO A 33	7 75.740	29.069	24.761	1.00	35.26
	MOTA	3171	СВ	PRO A 33	7 76.156	30.449	25.234	1.00	35.76
35	MOTA	3172	CG	PRO A 33	7 75.564	31.327	24.192	1.00	36.30
33	MOTA	3173	С	PRO A 33	7 74.325	28.751	25.198	1.00	36.60
	MOTA	3174	0	PRO A 33	7 73.358	29.231	24.604	1.00	39.83
40	MOTA	3175	N	ALA A 33	8 , 74.218	27.909	26.218	1.00	36.08
	MOTA	3177	CA	ALA A 33	8 72.944	27.512	26.802	1.00	35.56
45	MOTA	3178	CB	ALA A 33	8 72.671	26.039	26.533	1.00	36.41
	MOTA	3179	С	ALA A 33	73.116	27.752	28.294	1.00	36.55
	ATOM	3180	0	ALA A 33	8 74.059	27.241	28.902	1.00	37.65
50	ATOM	3181	N	VAL A 33	9 72.256	28.576	28.878	1.00	36.13
	MOTA	3183	CA	VAL A 33	72.360	28.872	30.295	1.00	34.95
55	MOTA	3184	СВ	VAL A 33	9 72.771	30.346	30.538	1.00	33.81
	ATOM	3185	CG1	VAL A 33	74.046	30.667	29.776	1.00	36.08
	ATOM	3186	CG2	VAL A 33	71.658	31.293	30.139	1.00	33.93
60	ATOM	3187	С	VAL A 33	71.07	28.578	31.043	1.00	35.50
	ATOM	3188	0	VAL A 33	69.990	28.553	30.451	1.00	38.45
65	MOTA	3189	N	ASP A 34	71.208	28.295	32.335	1.00	33.66
05	MOTA	3191	CA	ASP A 34	10 70.078	3 28.029	33.213	1.00	31.12

	MOTA	3192	CB	ASP A	340	70.318	26.785	34.067	1.00	31.54
	MOTA	3193	CG	ASP A	340	69.813	25.520	33.411	1.00	33.33
5	ATOM	3194	OD1	ASP A	340	68.895	25.606	32.569	1.00	35.59
	MOTA	3195	OD2	ASP A	340	70.324	24.432	33.742	1.00	35.60
10	MOTA	3196	С	ASP A	340	70.003	29.235	34.108	1.00	29.12
10	MOTA	3197	0	ASP A	340	70.968	29.555	34.794	1.00	29.84
	MOTA	3198	N	VAL A	341	68.890	29.948	34.040	1.00	28.41
15	MOTA	3200	CA	VAL A	341	68.687	31.133	34.846	1.00	28.82
	ATOM	3201	СВ	VAL A	341	68.249	32.316	33.969	1.00	29.73
20	ATOM	3202	CG1	VAL A	341	67.868	33.496	34.831	1.00	27.89
20	MOTA	3203	CG2	VAL A	341	69.364	32.703	33.017	1.00	28.62
•	MOTA	3204	С	VAL A	341	67.592	30.833	35.852	1.00	30.90
25	MOTA	3205	0	VAL A	341	66.456	30.596	35.459	1.00	34.21
	MOTA	3206	N	GLN A	342	67.931	30.821	37.139	1.00	31.91
30	MOTA	3208	CA	GLN A	342	66.948	30.546	38.191	1.00	31.19
	MOTA	3209	CB	GLN A	342	67.375	29.345	39.039	1.00	31.37
	MOTA	3210	CG	GLN A	342	66.319	28.867	40.028	1.00	31.56
35	MOTA	3211	CD	GLN A	342	66.706	27.570	40.706	1.00	33.87
	MOTA	3212	OE1	GLN A	342	67.841	27.111	40.590	1.00	37.83
40	MOTA	3213	NE2	GLN A	342	65.770	26.979	41.431	1.00	35.30
	MOTA	3216	С	GLN A	342	66.752	31.761	39.088	1.00	31.20
	MOTA	3217	0	GLN A	342	67.724	32.374	39.537	1.00	31.67
45	MOTA	3218	N	ALA A	343	65.494	32.114	39.327	1.00	30.96
	MOTA	3220	CA	ALA A	343	65.169	33.254	40.169	1.00	28.72
50	MOTA	3221	CB	ALA A	343	64.042	34.054	39.563	1.00	28.77
	MOTA	3222	С	ALA A	343	64.782	32.778	41.552	1.00	28.72
	MOTA	3223	0	ALA A	343	64.011	31.826	41.700	1.00	30.49
55	ATOM	3224	N	PHE A	344	65.320	33.447	42.559	1.00	26.88
	MOTA	3226	CA	PHE A	344	65.040	33.125	43.945	1.00	26.71
60	ATOM	3227	CB	PHE A	344	66.316	32.718	44.673	1.00	21.35
	MOTA	3228	CG	PHE A	344	66.891	31.428	44.210	1.00	19.20
_	MOTA	3229	CD1	PHE A	344	67.806	31.396	43.173	1.00	20.33
65	MOTA	3230		PHE A		66.547	30.247	44.834	1.00	15.57
	MOTA	3231	CE1	PHE A	344	68.372	30.198	42.766	1.00	19.57

	ATOM	3232	CE2	PHE A 344	67.106	29.049	44.438	1.00 17.79
_	ATOM	3233	CZ	PHE A 344	68.023	29.026	43.401	1.00 19.23
5	MOTA	3234	С	PHE A 344	64.523	34.382	44.608	1.00 29.77
	MOTA	3235	0	PHE A 344	64.847	35.495	44.189	1.00 33.20
10	ATOM	3236	N	ALA A 345	63.712	34.206	45.637	1.00 29.97
	ATOM	3238	CA	ALA A 345	63.204	35.331	46.390	1.00 30.90
	MOTA	3239	CB	ALA A 345	61.720	35.147	46.685	1.00 32.55
15	MOTA	3240	С	ALA A 345	64.030	35.266	47.669	1.00 31.52
	MOTA	3241	0	ALA A 345	64.230	34.186	48.220	1.00 32.91
20	MOTA	3242	N	VAL A 346	64.588	36.395	48.082	1.00 32.21
	ATOM	3244	CA	VAL A 346	65.403	36.449	49.287	1.00 33.87
25	ATOM	3245	СВ	VAL A 346	66.490	37.559	49.176	1.00 34.53
25	ATOM	3246	CG1	VAL A 346	67.430	37.537	50.383	1.00 32.13
	ATOM	3247	CG2	VAL A 346	67.276	37.394	47.892	1.00 33.38
30	ATOM	3248	, C	VAL A 346	64.497	36.755	50.474	1.00 36.21
	ATOM	3249	0	VAL A 346	63.985	37.871	50.597	1.00 38.00
35	ATOM	3250	N	LEU A 347	64.278	35.759	51.327	1.00 37.36
33	ATOM	3252	CA	LEU A 347	63.445	35.928	52.516	1.00 39.35
	MOTA	3253	CB	LEU A 347	63.172	34.567	53.162	1.00 37.87
40	ATOM	3254	CG	LEU A 347	62.664	33.460	52.238	1.00 37.83
	MOTA	3255	CD1	LEU A 347	62.315	32.231	53.057	1.00 36.57
45	MOTA	3256	CD2	LEU A 347	61.461	33.945	51.462	1.00 39.01
45	MOTA	3257	С	LEU A 347	64.157	36.847	53.519	1.00 42.26
	ATOM	3258	0	LEU A 347	65.376	37.021	53.449	1.00 44.26
50	ATOM	3259	N	PRO A 348	63.418	37.410	54.495	1.00 43.97
	ATOM	3260	CD	PRO A 348	61.973	37.254	54.728	1.00 43.89
55	ATOM	3261	CA	PRO A 348	64.002	38.309	55.505	1.00 43.61
J J	ATOM	3262	CB	PRO A 348	62.796	38.689	56.364	1.00 44.62
	MOTA	3263	CG	PRO A 348	61.875	37.514	56.206	1.00 44.98
60	ATOM	3264	С	PRO A 348	65.138	37.751	56.364	1.00 42.78
	MOTA	3265	0	PRO A 348	65.685	38.471	57.195	1.00 43.70
65	MOTA	3266	N	ASN A 349	65.478	36.478	56.184	1.00 42.71
<i>33</i>	MOTA	3268	CA	ASN A 349	66.556	35.849	56.947	1.00 41.55

	MOTA	3269	СВ	ASN A	349	66.046	34.591	57.659	1.00 41.16
	ATOM.	3270	CG	ASN A	349	65.583	33.518	56.689	1.00 43.53
5	MOTA	3271	OD1	ASN A	349	65.524	33.748	55.479	1.00 44.53
	MOTA	3272	ND2	ASN A	349	65.256	32.339	57.210	1.00 43.58
10	ATOM	3275	С	ASN A	349	67.731	35.482	56.038	1.00 41.09
10	ATOM	3276	0	asn a	349	68.662	34.801	56.469	1.00 41.04
	ATOM	3277	N	SER A	350	67.678	35.954	54.793	1.00 40.23
15	ATOM	3279	CA	SER A	350	68.697	35.704	53.768	1.00 39.62
	ATOM	3280	СВ	SER A	350	70.122	35.854	54.314	1.00 40.62
20	MOTA	3281	OG	SER A	350	70.518	37.214	54.317	1.00 46.79
,	ATOM	3283	С	SER A	350	68.561	34.364	53.067	1.00 37.74
	ATOM	3284	0	SER A	350	69.339	34.063	52.168	1.00 40.17
25	MOTA	3285	N	ALA A	351	67.581	33.562	53.471	1.00 34.00
	MOTA	3287	CA	ALA A	351	67.373	32.266	52.846	1.00 31.70
30	MOTA	3288	СВ	ALA A	351	66.426	31.429	53.670	1.00 31.56
50	ATOM	3289	С	ALA A	351	66.799	32.505	51.455	1.00 32.27
	ATOM	3290	0	ALA A	351	66.150	33.526	51.212	1.00 34.50
35	MOTA	3291	N	LEU A	352	67.022	31.560	50.549	1.00 30.81
	ATOM	3293	CA	LEU A	352	66.541	31.681	49.179	1.00 27.71
40	ATOM	3294	CB	LEU A	352	67.678	31.358	48.203	1.00 22.61
	ATOM	3295	CG	LEU A	352 .	69.026	32.057	48.373	1.00 14.74
	ATOM	3296	CD1	LEU A	352	69.958	31.612	47.280	1.00 13.29
45	MOTA	3297	CD2	LEU A	352	68.853	33.554	48.306	1.00 14.68
	ATOM	3298	С	LEU A	352	65.348	30.775	48.886	1.00 29.37
50	MOTA	3299	0	LEU A	352	65.375	29.573	49.179	1.00 30.14
	MOTA	3300	N	ALA A	353	64.309	31.349	48.289	1.00 31.25
	MOTA	3302	CA	ALA A	353	63.110	30.592	47.929	1.00 34.26
55	ATOM	3303	CB	ALA A	353	61.882	31.226	48.554	1.00 35.51
	MOTA	3304	С	ALA A	353	62.982	30.555	46.403	1.00 36.03
60	ATOM	3305	0	ALA A	353	62.793	31.587	45.764	1.00 37.52
	MOTA	3306	N	SER A	354	63.110	29.362	45.832	1.00 37.41
	ATOM	3308	CA	SER A	354	63.054	29.162	44.388	1.00 37.85
65	ATOM	3309	СВ	SER A	354	63.338	27.68 7	44.070	1.00 40.08
	ATOM	3310	OG	SER A	354	63.380	27.434	42.673	1.00 45.78

	MOTA	3312	Ċ	SER A	A	354	61.744	29.601	43.741	1.00 37.70
5	MOTA	3313	0	SER A	A	354	60.684	29.060	44.043	1.00 40.21
3	MOTA	3314	N	LEU 2	A	355	61.830	30.572	42.840	1.00 36.34
	MOTA	3316	CA	LEU 2	A	355	60.657	31.074	42.137	1.00 35.04
10	MOTA	3317	СВ	LEU 2	A	355	60.839	32.553	41.782	1.00 30.99
	ATOM	3318	CG	LEU .	A	355	60.830	33.547	42.939	1.00 28.58
1.5	ATOM	3319	CD1	LEU 2	A	355	61.028	34.955	42.423	1.00 28.81
15	MOTA	3320	CD2	LEU 2	A	355	59.520	33.441	43.678	1.00 30.20
	ATOM	3321	С	LEU .	A	355	60.399	30.255	40.868	1.00 36.96
20	MOTA	3322	0	LEU 2	A	355	59.363	29.592	40.745	1.00 36.42
	MOTA	3323	N	PHE .	A	356	61.356	30.286	39.939	1.00 37.24
0.5	ATOM	3325	CA	PHE .	Α	356	61.251	29.556	38.673	1.00 36.22
25	MOTA	3326	СВ	PHE .	A	356	60.315	30.291	37.699	1.00 33.82
	MOTA	3327	CG	PHE	Α	356	60.661	31.741	37.491	1.00 30.90
30	ATOM	3328	CD1	PHE .	A	356	61.786	32.110	36.768	1.00 29.04
	ATOM	3329	CD2	PHE .	Α	356	59.859	32.739	38.027	1.00 30.48
35	ATOM	3330	CE1	PHE .	A	356	62.111	33.451	36.586	1.00 28.29
33	ATOM	3331	CE2	PHE	A	356	60.179	34.081	37.848	1.00 30.44
	MOTA	3332	CZ	PHE .	A	356	61.308	34.435	37.128	1.00 25.36
40	MOTA	3333	С	PHE	A	356	62.611	29.356	38.011	1.00 36.47
	MOTA	3334	0	PHE	A	356	63.546	30.112	38.271	1.00 39.86
45	MOTA	3335	N	LEU	A	357	62.707	28.345	37.151	1.00 36.72
43	MOTA	3337	CA	LEU	A	357	63.937	28.035	36.423	1.00 35.97
	ATOM	3338	СВ	LEU	A	357	64.358	26.585	36.670	1.00 34.25
50	MOTA	3339	CG	LEU	A	357	65.593	26.104	35.904	1.00 32.69
	MOTA	3340	CD1	LEU	A	357	66.828	26.854	36.354	1.00 30.30
55	MOTA	3341	CD2	LEU	A	357	65.784	24.620	36.120	1.00 34.02
<i></i>	MOTA	3342	С	LEU	A	357	63.721	28.252	34.925	1.00 36.64
	MOTA	3343	0	LEU	A	357	62.805	27.678	34.333	1.00 39.03
60	MOTA	3344	N	ILE	A	358	64.585	29.054	34.318	1.00 35.88
	MOTA	3346	CA	ILE	A	358	64.508	29.367	32.899	1.00 34.57
65	MOTA	3347	CB	ILE	A	358	64.510	30.903	32.672	1.00 33.20
	MOTA	3348	CG2	ILE	A	358	64.653	31.239	31.208	1.00 31.65

	ATOM	3349	CG1	ILE A	358	63.220	31.510	33.207	1.00	33.82
	ATOM	3350	CD1	ILE A	358	61.978	30.907	32.602	1.00	34.80
5	ATOM	3351	С	ILE A	358	65.677	28.748	32.143	1.00	36.24
	ATOM	3352	0	ILE A	358	66.784	28.628	32.666	1.00	36.67
10	ATOM	3353	N	GLY A	359	65.405	28.338	30.912	1.00	38.02
10	ATOM	3355	CA	GLY A	359	66.419	27.753	30.061	1.00	37.56
	ATOM	3356	С	GLY A	359	66.597	28.704	28.902	1.00	38.10
15	ATOM	3357	0	GLY A	359	65.679	28.932	28.113	1.00	38.54
	ATOM	3358	N	MET A	360	67.769	29.305	28.830	1.00	37.31
20	MOTA	3360	CA	MET A	360	68.060	30.252	27.785	1.00	36.92
20	ATOM	3361	CB	MET A	360	68.515	31.560	28.405	1.00	38.52
	ATOM	3362	CG	MET A	360	68.813	32.656	27.421	1.00	42.50
25	MOTA	3363	SD	MET A	360	69.679	33.987	28.246	1.00	49.37
	ATOM	3364	CE	MET A	360	68.747	34.125	29.725	1.00	46.11
30	MOTA	3365	С	MET A	360	69.154	29.686	26.907	1.00	38.15
50	MOTA	3366	0	MET A	360	69.986	28.899	27.347	1.00	38.64
	ATOM	3367	N	HIS A	361	69.114	30.054	25.641	1.00	39.81
35	MOTA	3369	CA	HIS A	361	70.095	29.612	24.667	1.00	41.81
	ATOM	3370	CB	HIS A	361	69.766	28.215	24.131	1.00	45.81
40	MOTA	3371	CG	HIS A	361	68.310	27.988	23.878	1.00	51.49
	ATOM	3372	CD2	HIS A	361	67.290	27.716	24.728	1.00	54.71
	ATOM	3373	ND1	HIS A	361	67.753	28.041	22.618	1.00	53.69
45	ATOM	3375	CE1	HIS A	361	66.455	27.813	22.702	1.00	55.98
	ATOM	3376	NE2	HIS A	361	66.148	27.614	23.972	1.00	57.77
50	ATOM	3378	С	HIS A	361	70.032	30.642	23.569	1.00	41.72
	ATOM	3379	0	HIS A	361	69.003	31.295	23.387	1.00	42.60
	ATOM	3380	N	THR A	362	71.132	30.816	22.858	1.00	40.54
55	ATOM	3382	CA	THIR A	362	71.168	31.803	21.803	1.00	38.28
	ATOM	3383	CB	THR A	362	71.452	33.206	22.399	1.00	38.27
60	ATOM	3384	OG1	THR A	362	71.408	34.200	21.371	1.00	40.01
	ATOM	3386	CG2	THR A	362	72.799	33.241	23.082	1.00	38.36
	ATOM	3387	С	THR A	362	72.244	31.391	20.818	1.00	37.84
65	MOTA	3388	0	THR A	362	72.692	30.245	20.829	1.00	38.47
	MOTA	3389	N	THR A	363	72.620	32.315	19.948	1.00	37.83

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	MOTA	3391	CA	THR A 363	73.640	32.091	18.940	1.00 38.05
5	ATOM	3392	СВ	THR A 363	73.007	31.617	17.603	1.00 38.92
3	ATOM	3393	OG1	THR A 363	73.993	31.618	16.567	1.00 41.55
	ATOM	3395	CG2	THR A 363	71.855	32.515	17.193	1.00 42.85
10	ATOM	3396	С	THR A 363	74.333	33.436	18.773	1.00 36.99
	ATOM	3397	0	THR A 363	73.668	34.467	18.705	1.00 38.50
1.5	ATOM	3398	N	GLY A 364	75.660	33.438	18.763	1.00 36.22
15	ATOM	3400	CA	GLY A 364	76.377	34.692	18.626	1.00 35.40
	ATOM	3401	С	GLY A 364	77.639	34.632	17.794	1.00 35.46
20	ATOM	3402	0	GLY A 364	78.075	33.561	17.395	1.00 35.68
	ATOM	3403	N	SER A 365	78.216	35.795	17.522	1.00 37.37
25	ATOM	3405	CA	SER A 365	79.434	35.884	16.738	1.00.39.81
23	MOTA	3406	CB	SER A 365	79.399	37.121	15.840	1.00 41.52
	MOTA	3407	OG	SER A 365	78.345	37.028	14.895	1.00 50.49
30	MOTA	3409	С	SER A 365	80.627	35.975	17.669	1.00 40.48
	MOTA	3410	0	SER A 365	80.584	36.669	18.682	1.00 43.48
35	MOTA	3411	N	MET A 366	81.685	35.259	17.330	1.00 40.04
55	MOTA	3413	CA	MET A 366	82.898	35.271	18.118	1.00 40.25
	MOTA	3414	CB	MET A 366	83.254	33.836	18.509	1.00 41.64
40	MOTA	3415	CG	MET A 366	84.524	33.688	19.310	1.00 44.41
	MOTA	3416	SD	MET A 366	84.378	34.496	20.881	1.00 48.93
45	ATOM	3417	CE	MET A 366	83.512	33.252	21.833	1.00 49.58
	MOTA	3418	С	MET A 366	83.978	35.880	17.224	1.00 41.24
	MOTA	3419	0	MET A 366	84.400	35.262	16.249	1.00 43.55
50	ATOM	3420	N	GLU A 367	84.347	37.129	17.482	1.00 42.13
	MOTA	3422	CA	GLU A 367	85.383	37.780	16.686	1.00 43.88
55	MOTA	3423	CB	GLU A 367	85.148	39.288	16.608	1.00 47.79
	MOTA	3424	CG	GLU A 367	84.094	39.715	15.588	1.00 54.26
	MOTA	3425	CD	GLU A 367	84.551	39.555	14.143	1.00 57.86
60	MOTA	3426	OEI	L GLU A 367	85.768	39.681	13.872	1.00 60.11
	MOTA	3427	OE2	2 GLU A 367	83.683	39.315	13.273	1.00 59.61
65	MOTA	3428	С	GLU A 367	86.724	37.492	17.336	1.00 43.57
	ATOM	3429	0	GLU A 367	86.822	37.491	18.562	1.00 44.51

	MOTA	3430	N	VAL A	368	87.752	37.256	16.526	1.00	42.06
	MOTA	3432	CA	VAL A	368	89.077	36.944	17.047	1.00	42.38
5	MOTA	3433	CB	VAL A	368	89.412	35.454	16.810	1.00	42.31
	MOTA	3434	CG1	VAL A	368	90.866	35.167	17.146	1.00	44.40
10	ATOM	3435	CG2	VAL A	A 368	88.509	34.581	17.660	1.00	41.44
10	ATOM	3436	С	VAL A	368	90.204	37.822	16.491	1.00	43.82
	ATOM	3437	0	VAL A	368	90.306	38.039	15.279	1.00	44.42
15	ATOM	3438	N	SER F	369	91.051	38.312	17.392	1.00	44.15
•	ATOM	3440	CA	SER A	369	92.182	39.160	17.032	1.00	46.20
20	ATOM	3441	СВ	SER A	A 369	91.838	40.636	17.254	1.00	48.08
20	ATOM	3442	OG	SER A	A 369	90.692	41.011	16.508	1.00	55.65
	ATOM	3444	С	SER A	A 369	93.376	38.773	17.896	1.00	46.39
25	ATOM	3445	0	SER A	A 369	93.269	37.897	18.757	1.00	45.59
	ATOM	3446	N	ALA A	A 370	94.503	39.447	17.692	1.00	46.96
30	ATOM	3448	CA	ALA A	A 370	95.705	39.151	18.455	1.00	46.86
50	MOTA	3449	CB	ALA A	A 370	96.659	38.334	17.615	1.00	48.51
	ATOM	3450	С	ALA A	A 370	96.401	40.399	18.961	1.00	47.76
35	MOTA	3451	0	ALA A	A 370	96.526	41.384	18.234	1.00	49.78
	MOTA	3452	N	GLU 1	A 371	96.833	40.346	20.215	1.00	49.05
40	MOTA	3454	CA	GLU A	A 371	97.546	41.429	20.889	1.00	52.58
-	MOTA	3455	CB	GLU Z	A 371	96.670	42.035	21.999	1.00	.55.47
	MOTA	3456	CG	GLU Z	A 371	97.394	42.954	23.010	1.00	61.82
45	MOTA	3457	CD	GLU .	A 371	97.662	44.365	22.494	1.00	66.05
	ATOM	3458	OE1	GLU	A 371	96.728	44.994	21.945	1.00	69.54
50	ATOM	3459	OE2	GLU .	A 371	98.801	44.859	22.662	1.00	66.75
	ATOM	3460	С	GLU .	A 371	98.753	40.718	21.490	1.00	53.47
	ATOM	3461	0	GLU .	A 371	98.617	39.613	22.015	1.00	54.95
55	MOTA	3462	N	SER	A 372	99.926	41.337	21.418	1.00	54.76
	ATOM	3464	CA	SER	A 372	101.145	40.718	21.932	1.00	54.93
60	MOTA	3465	СВ	SER	A 372	101.080	40.570	23.462	1.00	56.19
	MOTA	3466	OG	SER	A 372	100.788	41.803	24.098	1.00	57.76
	MOTA	3468	С	SER	A 372	101.267	39.347	21.245	1.00	54.65
65	MOTA	3469	0	SER	A 372	101.364	39.275	20.017	1.00	56.14
	MOTA	3470	N	ASN	A 373	101.207	38.271	22.023	1.00	51.39

	MOTA	3472	CA	ASN A	373	101.292	36.922	21.480	1.00	48.43
_	MOTA	3473	CB	ASN A	373	102.606	36.249	21.891	1.00	49.96
5	MOTA	3474	CG	ASN A	373	103.830	36.981	21.379	1.00	51.10
	ATOM	3475	OD1	ASN A	373	103.780	37.674	20.361	1.00	53.30
10	ATOM .	3476	ND2	ASN A	373	104.946	36.818	22.078	1.00	50.08
	MOTA	3479	С	ASN A	373	100.126	36.168	22.087	1.00	46.52
16	MOTA	3480	0	ASN A	373	100.267	35.027	22.529	1.00	45.55
15	MOTA	3481	N	ARG A	374	98.983	36.838	22.156	1.00	44.90
	MOTA	3483	CA	ARG A	374	97.780	36.255	22.730	1.00	43.78
20	ATOM	3484	СВ	ARG A	374	97.293	37.082	23.929	1.00	45.07
	MOTA	3485	CG	ARG A	A 374	98.263	37.183	25.090	1.00	47.75
25	MOTA	3486	CD	ARG A	374	97.653	37.976	26.235	1.00	52.92
23	MOTA	3487	NE	ARG A	374	97.293	39.342	25.857	1.00	58.67
	MOTA	3489	CZ	ARG A	374	97 . 7 37	40.439	26.468	1.00	62.31
30	MOTA	3490	NHl	ARG A	374	98.568	40.347	27.500	1.00	63.80
	MOTA	3493	NH2	ARG A	374	97.345	41.637	26.047	1.00	62.63
35	MOTA	3496	С	ARG A	A 374	96.660	36.163	21.706	1.00	41.67
33	MOTA	3497	0	ARG A	A 374	96.660	36.866	20.698	1.00	42.40
	ATOM	3498	N	LEU A	A 375	95.720	35.268	21.972	1.00	39.53
40	MOTA	3500	CA	LEU A	A 375	94.568	35.063	21.119	1.00	38.25
	ATOM	3501	CB	LEU A	A 375	94.366	33.568	20.886	1.00	40.19
45	MOTA	3502	CG	LEU A	A 375	93.330	33.154	19.846	1.00	42.29
	ATOM	3503	CD1	LEU A	A 375	93.784	33.625	18.478	1.00	42.24
	ATOM	3504	CD2	TEO 1	A 375	93.158	31.646	19.866	1.00	41.82
50	MOTA	3505	C	LEU 2	A 375	93.399	35.623	21.917		37.82
	MOTA	3506	0	LEU A	A 375	93.024	35.058	22.946	1.00	38.21
55	MOTA	3507	N		A 376	92.871	36.764	21.491	1.00	38.16
	MOTA	3509	CA		A 376	91.752	37.384			
	ATOM	3510	CB		A 376	92.057				37.73
60	ATOM	3511		VAL 2		93.386	38.928			36.49
	MOTA	3512		VAL .		92.054				37.89
65	ATOM	3513	С		A 376	90.517	37.376	21.313	1.00	37.85
	MOTA	3514	0	VAL .	A 376	90.615	37.251	20.091	1.00	37.84

	MOTA	3515	N	GLY	A	377	89.358	37.542	21.935	1.00	38.79
	ATOM	3517	CA	GLY	A	377	88.120	37.548	21.186	1.00	38.96
5	ATOM	3518	С	GLY	A	377	87.034	38.334	21.880	1.00	38.86
	MOTA	3519	0	GLY	A	377	87.235	38.860	22.976	1.00	40.46
10	ATOM	3520	N	GLU	A	378	85.881	38.426	21.235	1.00	39.21
10	MOTA	3522	CA	GLU	A	378	84.752	39.143	21.793	1.00	39.89
	ATOM	3523	СВ	GLU	A	378	84.762	40.601	21.335	1.00	42.06
15	MOTA	3524	CG	GLU	A	378	83.935	41.522	22.226	1.00	47.89
	MOTA	3525	CD	GLU	A	378	83.384	42.725	21.488	1.00	51.11
20	MOTA	3526	OE1	GLU	A	378	84.142	43.357	20.718	1.00	53.99
20	ATOM	3527	OE2	GLU	A	378	82.184	43.031	21.679	1.00	52.17
	MOTA	3528	С	GLU	A	378	83.481	38.456	21.315	1.00	40.11
25	ATOM	3529	0	GLU	A	378	83.352	38.132	20.133	1.00	40.78
	MOTA	3530	N	LEU	A	379	82.569	38.211	22.249	1.00	40.26
30	MOTA	3532	CA	LEU	A	379	81.297	37.558	21.973	1.00	41.54
50	ATOM	3533	СВ	LEU	A	379	80.935	36.651	23.155	1.00	43.08
	ATOM	3534	CG	LEU	A	379	80.208	35.314	22.991	1.00	45.34
35	MOTA	3535	CD1	LEU	A	379	79.839	34.798	24.377	1.00	44.87
	ATOM	3536	CD2	LEU	A	379	78.966	35.444	22.136	1.00	44.17
40	ATOM	3537	С	LEU	A	379	80.199	38.613	21.799	1.00	42.68
	MOTA	3538	0	LEU	A	379	80.179	39.623	22.507	1.00	43.44
	ATOM	3539	N	LYS	A	380	79.303	38.384	20.845	1.00	44.09
45	ATOM	3541	CA	LYS	A	380	78.171	39.276	20.588	1.00	46.18
	MOTA	3542	CB	LYS	A	380	78.438	40.165	19.373	1.00	48.46
50	MOTA	3543	CG	LYS	A	380	79.444	41.279	19.660	1.00	54.25
	MOTA	3544	CD	LYS	A	380	79.847	42.041	18.408	1.00	58.33
	MOTA	3545	CE	LYS	A	380	80.817	43.172	18.734	1.00	60.62
55	MOTA	3546	NZ			380	81.332	43.851	17.509	1.00	63.04
	MOTA	3550	С			380	76.968	38.367	20.369		46.00
60	MOTA	3551	0	LYS	A	380	77.042	37.418	19.591	1.00	45.80
	MOTA	3552	.N	LEU	A	381	75.877	38.630	21.082	1.00	46.45
4.5	MOTA	3554	CA			381	74.690	37.784	20.996		46.82
65 ·	MOTA	3555	CB			381	74.115		22.400	1.00	45.07
	MOTA	3556	CG	LEU	A	381	75.001	36.992	23.521	1.00	41.15

	MOTA	3557	CD1	LEU A	4	381	74.206	36.974	24.817	1.00	39.28
~	MOTA	3558	CD2	LEU A	Ą	381	75.500	35.604	23.174	1.00	40.04
5	MOTA	3559	С	LEU A	Ą	381	73.569	38.272	20.084	1.00	48.41
	MOTA	3560	0	LEU A	A	381	73.505	39.448	19.719	1.00	48.91
10	MOTA	3561	N	ASP A	4	382	72.696	37.335	19.725	1.00	50.06
	ATOM	3563	CA	ASP A	Ą	382	71.527	37.591	18.894	1.00	54.68
ic	MOTA	3564	СВ	ASP A	Ą	382	71.371	36.488	17.845	1.00	57.78
15	ATOM	3565	CG	ASP A	A	382	71.718	36.953	16.453	1.00	61.73
	MOTA	3566	OD1	ASP A	Ą	382	72.846	37.455	16.258	1.00	65.65
20	ATOM	3567	OD2	ASP A	A	382	70.862	36.809	15.552	1.00	62.89
	ATOM	3568	С	ASP A	A	382	70.325	37.565	19.836	1.00	56.30
25	ATOM	3569	0	ASP A	A	382	70.465	37.827	21.030	1.00	57.37
25	ATOM	3570	N	ARG A	A	383	69.147	37.244	19.304	1.00	57.17
	ATOM	3572	CA	ARG 2	A	383	67.932	37.169	20.116	1.00	56.86
30	ATOM	3573	СВ	ARG .	A	383	66.710	36.913	19.221	1.00	61.20
	ATOM	3574	CG	ARG .	A.	383	66.368	38.054	18.262	1.00	66.61
35	MOTA	3575	CD	ARG .	A	383	65.158	37.721	17.389	1.00	68.98
33	MOTA	3576	NE	ARG	Α	383	64.674	38.887	16.647	1.00	73.50
	ATOM	3578	CZ	ARG	A	383	64.914	39.119	15.358	1.00	76.60
40	ATOM	3579	NH1	ARG	A	383	65.639	38.269	14.636	1.00	77.94
	MOTA	3582	NH2	ARG	A	383	64.421	40.210	14.784	1.00	77.33
45	ATOM	3585	С	ARG	A	383	68.081	36.026	21.120	1.00	54.61
43	MOTA	3586	0	ARG	A	383	68.689	35.001	20.801	1.00	56.02
	MOTA	3587	N	LEU	A	384	67.565	36.207	22.333	1.00	50.19
50	MOTA	3589	CA	LEU	A	384	67.654	35.163	23.353	1.00	47.21
	MOTA	3590	СВ	LEU	A	384	67.886	35 .765	24.740	1.00	45.34
55	MOTA	3591	CG	LEU	A	384	69.072	36.707	24.951	1.00	44.13
55	MOTA	3592	CD1	LEU	A	384	69.089	37.136	26.403	1.00	48.33
	MOTA	3593	CD2	LEU	A	384	70.386	36.036	24.587	1.00	44.37
60	ATOM	3594	С	LEU	A	384	66.366	34.355	23.360	1.00	46.05
	ATOM	3595	0	LEU	A	384	65.287	34.910	23.175	1.00	46.98
65	MOTA	3596	N	LEU	A	385	66.477	33.044	23.536	1.00	45.63
UJ	ATOM	3598	CA	LEU	A	385	65.299	32.190	23.564	1.00	44.83

	MOTA	3599	СВ	LEU	A	385	65.417	31.034	22.577	1.00	47.07
	ATOM	3600	CG	LEU	A	385	65.424	31.305	21.075	1.00	49.68
5	MOTA	3601	CD1	LEU	A	385	66.807	31.742	20.610	1.00	50.06
	MOTA	3602	CD2	LEU	A	385	65.018	30.020	20.367	1.00	53.29
10	ATOM	3603	С	LEU	A	385	65.093	31.624	24.948	1.00	43.70
10	ATOM	3604	0	LEU	A	385	65.973	30.965	25.500	1.00	46.13
	ATOM	3605	Ŋ	LEU	A	386	63.921	31.879	25.501	1.00	41.90
15	ATOM	3607	CA	LEU	A	386	63.586	31.399	26.819	1.00	40.93
	ATOM	3608	СВ	LEU	À	386	62.804	32.467	27.576	1.00	38.95
20	ATOM	3609	CG	LEU	A	386	63.610	33.573	28.251	1.00	38.87
20	ATOM	3610	CD1	LEU	A	386	64.732	34.076	27.364	1.00	36.69
	ATOM	3611	CD2	LEU	A	386	62.671	34.690	28.636	1.00	38.24
25	MOTA	3612	С	LEU	A	386	62.754	30.140	26.711	1.00	41.89
	MOTA	3613	0	LEU	A	386	62.154	29.861	25.677	1.00	44.73
30	MOTA	3614	N	GLU	A	387	62.777	29.354	27.773	1.00	42.45
30	MOTA	3616	CA	GLU	A	387	62.006	28.133	27.867	1.00	43.22
	ATOM	3617	СВ	GLU	A	387	62.773	26.932	27.306	1.00	44.48
35	ATOM	3618	CĢ	GLU	A	387	61.993	25.614	27.408	1.00	46.64
	ATOM	3619	CD	GLU	A	387	62.699	24.436	26.762	1.00	48.74
40	ATOM	3620	OE1	GLU	A	387	62.622	24.305	25.520	1.00	53.50
, ,	ATOM	3621	OE2	GLU	A	387	63.311	23.629	27.495	1.00	48.42
	ATOM	3622	С	GLU	A	387	61.791	27.966	29.359	1.00	45.59
45	ATOM	3623	0	GLU	A	387	62.697	28.232	30.150	1.00	47.68
	ATOM	3624	N	LEU	A	388	60.574	27.618	29.750	1.00	47.18
50	MOTA	3626	CA	LEU	A	388	60.260	27.422	31.156	1.00	48.51
	MOTA	3627	CB	LEU	A	388	58.795	27.778	31.418	1.00	49.74
	ATOM	3628	CG	LEU	A	388	58.298	27.776	32.865	1.00	49.97
55	MOTA	3629	CD1	LEU	A	388	58.957	28.902	33.656	1.00	49.70
	MOTA	3630	CD2	LEU	A	388	56.788	27.935	32.871	1.00	51.43
60	MOTA	3631	С	LEU	A	388	60.513	25.953	31.468	1.00	49.89
	ATOM	3632	0	LEU	A	388	60.033	25.071	30.754	1.00	49.90
	MOTA	3633	N	LYS	A	389	61.290	25.688	32.511	1.00	53.07
65	MOTA	3635	CA	LYS	A	389	61.599	24.312	32.883	1.00	57.61
	MOTA	3636	CB	LYS	A	389	63.109	24.152	33.105	1.00	58.15

	ATOM	3637	CG	LYS A	389	63.933	24.486	31.864	1.00 58.89
5	MOTA	3638	CD	LYS A	389 ·	65.377	24.022	31.971	1.00 61.45
3	ATOM	3639	CE	LYS A	389	66.104	24.202	30.640	1.00 63.53
	ATOM	3640	NZ	LYS A	389	67.507	23.692	30.664	1.00 66.07
10	ATOM	3644	С	LYS A	389	60.800	23.813	34.094	1.00 60.00
	MOTA	3645	0	LYS A	389	60.351	22.663	34.122	1.00 61.33
1.5	MOTA	3646	N	HIS A	390	60.622	24.679	35.086	1.00 61.51
15	MOTA	3648	CA	HIS A	390	59.871	24.347	36.297	1.00 63.22
	ATOM	3649	СВ	HIS A	390	60.713	23.474	37.242	1.00 68.01
20	MOTA	3650	CG	HIS A	390	59.939	22.880	38.386	1.00 74.01
	ATOM	3651	CD2	HIS A	390	59.403	21.647	38.553	1.00 76.41
25	ATOM	3652	ND1	HIS A	390	59.682	23.568	39.554	1.00 76.75
25	MOTA	3654	CE1	HIS A	390	59.023	22.785	40.391	1.00 78.97
	MOTA	3655	NE2	HIS A	390	58.843	21.613	39.807	1.00 79.75
30	ATOM	3657	С	HIS A	390	59.551	25.669	36.969	1.00 61.81
	ATOM	3658	0	HIS A	390	60.348	26.604	36.912	1.00 62.27
35	MOTA	3659	N	SER A	391	58.375	25.762	37.572	1.00 60.92
55	MOTA	3661	CA	SER A	391	57.978	26.980	38.257	1.00 59.76
	MOTA	3662	СВ	SER A	391	57.005	27.793	37.403	1.00 58.20
40	MOTA	3663	OG	SER A	391 .	56.639	28.996	38.059	1.00 55.50
	MOTA	3665	С	SER A	391	57.322	26.605	39.567	1.00 59.99
45	MOTA	3666	0	SER A	391	56.486	25.699	39.618	1.00 60.83
43	ATOM	3667	Ŋ	ASN A	392	57.753	27.260	40.636	1.00 59.65
	MOTA	3669	CA	ASN A	392	57.190	27.008	41.949	1.00 59.99
50	MOTA	3670	СВ	ASN A	392	58.241	27.234	43.039	1.00 58.88
	MOTA	3671	CG	ASN A	392	59.314	26.152	43.054	1.00 58.15
55	MOTA	3672	OD1	ASN A	392	59.424	25.344	42.126	1.00 59.13
<i>33</i>	MOTA	3673	ND2	ASN A	392	60.117	26.137	44.108	1.00 57.91
	MOTA	3676	С	ASN A	392	55.976	27.914	42.145	1.00 60.50
60	MOTA	3677	0	asn a	392	55.149	27.678	43.023	1.00 61.62
	MOTA	3678	N	ILE A	393	55.851	28.926	41.288	1.00 61.07
65	ATOM	3680	CA	ILE A	393	54.730	29.861	41.351	1.00 60.89
V 5	ATOM	3681	СВ	ILE A	393	55.212	31.333	41.306	1.00 58.61

	MOTA	3682	CG2	ILE A	393	56.083	31.634	42.510	1.00	58.38
	MOTA	3683	CG1	ILE A	393	55.948	31.622	39.995	1.00	57.06
5	MOTA	3684	CD1	ILE A	393	56.245	33.087	39.775	1.00	56.51
	MOTA	3685	С	ILE A	393	53.700	29.632	40.228	1.00	63.24
10	ATOM	3686	0	ILE A	393	53.049	30.574	39.772	1.00	64.68
10	ATOM	3687	N	GLY A	394	53.556	28.382	39.786	1.00	64.29
	ATOM	3689	CA	GLY A	394	52.603	28.061	38.731	1.00	62.76
15	MOTA	3690	С	GLY A	394	53.046	28.426	37.323	1.00	62.37
	ATOM	3691	0	GLY A	394	53.985	29.202	37.147	1.00	62.54
20	MOTA	3692	N	PRO A	395	52.412	27.848	36.291	1.00	62.40
20	MOTA	3693	CD	PRO A	395	51.333	26.846	36.373	1.00	63.52
	ATOM	3694	CA	PRO A	395	52.758	28.130	34.894	1.00	60.73
25	MOTA	3695	СВ	PRO A	395	51.937	27.095	34.125	1.00	61.59
	MOTA	3696	CG	PRO A	395	50.728	26.910	34.992	1.00	63.08
30	MOTA	3697	С	PRO A	395	52.405	29.554	34.470	1.00	59.25
J 0	MOTA	3698	0	PRO A	395	51.353	30.081	34.834	1.00	59.47
	MOTA	3699	N	PHE A	396	53.295	30.171	33.702	1.00	57.50
35	ATOM	3701	CA	PHE A	396	53.087	31.530	33.220	1.00	55.80
	MOTA	3702	CB	PHE A	396	53.684	32.555	34.200	1.00	54.39
40	MOTA	3703	CG	PHE A	396	55.194	32.588	34.223	1.00	51.73
	MOTA	3704	CD1	PHE A	396	55.899	33.426	33.364	1.00	49.07
	MOTA	3705	CD2	PHE A	396	55.909	31.796	35.115	1.00	50.73
45	MOTA	3706	CE1	PHE A	396	57.283	33.474	33.394	1.00	47.56
	MOTA	3707	CE2	PHE A	396	57.295	31.841	35.150	1.00	47.47
50	ATOM	3708	CZ	PHE A	396	57.981	32.678	34.289	1.00	47.02
	ATOM	3709	C	PHE A	396	53.720	31.674	31.839	1.00	55.76
	MOTA	3710	0	PHE A	396	54.463	30.792	31.392	1.00	56.39
55	ATOM	3711	N	PRO A	397	53.394	32.763	31.120	1.00	54.61
	ATOM	3712	CD	PRO A	397	52.354	33.768	31.393	1.00	54.50
60	ATOM	3713	CA	PRO A	397	53.966	32.967	29.787	1.00	53.13
	ATOM	3714	СВ	PRO A	397	53.167	34.159	29.252	1.00	53.76
	ATOM	3715	CG	PRO A	397	51.877	34.095	30.011	1.00	53.30
65	ATOM	3716	С	PRO A	. 397	55.448	33.301	29.899	1.00	51.99
	ATOM	3717	0	PRO A	397	55.810	34.434	30.219	1.00	51.04

		ATOM	3718	N	VAL A	398	56.300	32.322	29.609	1.00 50.99
	_	MOTA	3720	CA	VAL A	398	57.747	32.506	29.686	1.00 50.88
	5	ATOM	3721	CB	VAL A	398	58.500	31.222	29.253	1.00 50.94
		ATOM	3722	CG1	VAL A	398	58.260	30.926	27.785	1.00 50.60
	10	ATOM	3723	CG2	VAL A	. 398	59.987	31.340	29.561	1.00 49.69
		ATOM	3724	С	VAL A	398	58.248	33.709	28.888	1.00 51.53
		ATOM	3725	0	VAL A	398	59.383	34.144	29.053	1.00 54.45
	15	MOTA	3726	N	GLU A	399	57.385	34.268	28.050	1.00 52.68
-		ATOM	3728	CA	GLU A	399	57.741	35.427	27.236	1.00 52.46
	20	MOTA	3729	СВ	GLU A	399	56.807	35.533	26.020	1.00 53.50
		MOTA	3730	CG	GLU A	399	56.637	34.234	25.210	1.00 53.56
		MOTA	3731	CD	GLU A	399	55.420	33.405	25.629	1.00 53.53
	25	ATOM	3732	OE1	GLU A	399	54.313	33.979	25.741	1.00 54.16
		ATOM	3733	OE2	GLU A	399	55.562	32.176	25.821	1.00 52.27
	30	MOTA	3734	С	GLU A	A 399	57.737	36.750	28.025	1.00 51.41
		ATOM	3735	0	GLU A	A 399	58.233	37.762	27.539	1.00 52.41
	25	ATOM	3736	N	LEU /	A 400	57.186	36.747	29.235	1.00 49.79
	35	MOTA	3738	CA	LEU A	4 400	57.148	37.955	30.060	1.00 49.44
		MOTA	3739	CB	LEU A	A 400	56.186	37.769	31.238	1.00 50.82
	40	MOTA	3740	CG	LEU :	A 400	54.683	38.030	31.086	1.00 49.58
		MOTA	3741	CD1	LEU	A 400	54.427	39.510	30.867	1.00 50.78
	45	MOTA	3742	CD2	LEU	A 400	54.114	37.220	29.954	1.00 50.81
	43	MOTA	3743	C	LEU :	A 400	58.539	38.330	30.589	1.00 50.26
		MOTA	3744	0	LEU .	A 400	58.800	39.487	30.931	1.00 50.71
	50	ATOM	3745	N	LEU	A 401	59.428	37.344	30.656	1.00 50.41
		ATOM	3747	CA	LEU	A 401	60.790	37.548	31.143	1.00 49.44
	55	MOTA	3748	CB	LEU	A 401	61.310	36.255	31.766	1.00 48.32
	JJ	ATOM	3749	CG	LEU	A 401	60.598	35.840	33.049	1.00 46.40
		MOTA	3750	CD:	r tea	A 401	60.738	34.361	33.262	1.00 45.30
	60	MOTA	3751	CD	LEU	A 401	61.168	36.620	34.218	1.00 48.02
		MOTA	3752	С	LEU	A 401	61.762	38.013	30.066	1.00 49.89
	65	ATOM	3753	0	LEU	A 401	62.919	38.303	30.360	1.00 51.08
	UJ	MOTA	3754	N	GLN	A 402	61.288	38.091	28.826	1.00 51.14

	MOTA	3756	CA	GLN A	402	62.107	38.511	27.686	1.00 52.43
	ATOM	3757	СВ	GLN A	402	61.255	38.569	26.413	1.00 53.25
5	ATOM	3758	CG	GLN A	402	60.908	37.211	25.810	1.00 55.42
	ATOM	3759	CD	GLN A	402	61.966	36.697	24.851	1.00 55.66
10	ATOM	3760	OE1	GLN A	402	63.075	37.229	24.780	1.00 55.93
10	ATOM	3761	NE2	GLN A	402	61.619	35.667	24.090	1.00 56.71
	ATOM	3764	С	GLN A	402	62.800	39.855	27.890	1.00 53.23
15	ATOM	3765	0	GLN A	402	64.009	39.977	27.684	1.00 54.78
	MOTA	3766	N	ASP A	403	62.031	40.866	28.275	1.00 53.89
	MOTA	3768	CA	ASP A	403	62.578	42.198	28.505	1.00 54.40
20	MOTA	3769	СВ	ASP A	403	61.441	43.183	28.797	1.00 58.20
	MOTA	3770	CG	ASP A	403	60.524	43.394	27.597	1.00 60.37
25	ATOM	3771	OD1	ASP A	403	61.001	43.915	26.563	1.00 60.48
	MOTA	3772	OD2	ASP A	403	59.325	43.045	27.691	1.00 61.61
30	ATOM	3773	С	ASP A	403	63.615	42.204	29.638	1.00 53.27
30	ATOM	3774	0	ASP A	403	64.669	42.836	29.525	1.00 53.59
	ATOM	3775	N	ILE A	404	63.318	41.487	30.719	1.00 51.11
35	ATOM	3777	CA	ILE A	404	64.229	41.387	31.852	1.00 48.30
	ATOM	3778	CB	ILE A	404	63.545	40.668	33.055	1.00 45.31
40	ATOM	3779	CG2	ILE A	404	64.352	39.468	33.530	1.00 47.52
40	ATOM	3780	CG1	ILE A	404	63.337	41.654	34.203	1.00 42.01
	ATOM	3781	CD1	ILE A	404	64.620	42.234	34.752	1.00 36.04
45	ATOM	3782	С	ILE A	404	65.497	40.656	31.400	1.00 48.48
	ATOM	3783	0	ILE A	404	66.611	41.108	31.667	1.00 51.31
50	ATOM	3784	N	MET A	405	65.318	39.564	30.660	1.00 47.51
	ATOM	3786	CA	MET A	405	66.433	38.768	30.151	1.00 47.38
	ATOM	3787	СВ	MET A	405	65.920	37.479	29.505	1.00 47.17
55	ATOM	3788	CG	MET A	405	66.151	36.233	30.345	1.00 48.66
	ATOM	3789	SD	MET A	405	65.913	36.487	32.115	1.00 49.07
60	ATOM	3790	CE	MET A	405	67.594	36.786	32.640	1.00 50.09
	ATOM	3791	C .	MET A	405	67.291	39.540	29.160	1.00 46.62
	ATOM	3792	0	MET A	405	68.500	39.339	29.092	1.00 48.06
65	ATOM	3793	N .	ASN A	406	66.666	40.441	28.413	1.00 46.76
	MOTA	3795	CA	ASN A	406	67.383	41.239	27.433	1.00 47.68

	MOTA	3796	CB	ASN .	A	406	66.467	41.675	26.295	1.00	51.33
5	MOTA	3797	CG	ASN .	A	406	66.534	40.732	25.115	1.00	54.24
3	MOTA	3798	OD1	ASN .	A	406	67.448	40.813	24.290	1.00	55.58
	ATOM	3799	ND2	ASN .	A	406	65.580	39.814	25.039	1.00	54.80
10	ATOM	3802	С	ASN .	A	406	68.134	42.430	27.999	1.00	46.60
	ATOM	3803	0	ASN .	A	406	68.803	43.148	27.263	1.00	48.59
1.5	MOTA	3804	N	TYR .	A	407	67.993	42.674	29.293	1.00	45.78
15	ATOM	3806	CA	TYR .	A	407	68.732	43.760	29.922	1.00	44.83
	ATOM	3807	СВ	TYR	A	407	67.825	44.613	30.829	1.00	46.29
20	ATOM	3808	CG	TYR	A	407	68.558	45.733	31.553	1.00	45.38
	MOTA	3809	CD1	TYR	A	407	68.891	46.916	30.897	1.00	45.96
25	MOTA	3810	CE1	TYR	A	407	69.621	47.915	31.540	1.00	47.49
23	ATOM	3811	CD2	TYR	A	407	68.966	45.582	32.880	1.00	45.49
	MOTA	3812	CE2	TYR	A	407	69.694	46.573	33.532	1.00	44.88
30	MOTA	3813	CZ	TYR	A	407	70.022	47.733	32.858	1.00	47.09
	MOTA	3814	ОН	TYR	A	407	70.771	48.701	33.493	1.00	48.41
35	MOTA	3816	С	TYR	A	407	69.837	43.080	30.730	1.00	43.05
33	MOTA	3817	0	TYR	A	407	71.024	43.262	30.462	1.00	42.68
	MOTA	3818	N	ILE	A	408	69.423	42.229	31.661	1.00	41.43
40	MOTA	3820	CA	ILE	A	408	70.332	41.490	32.523	1.00	40.32
	ATOM	3821	CB	ILE	A	408	69.580	40.371	33.272	1.00	40.57
45	ATOM	3822	CG2	ILE	A	408	70.550	39.445	33.978	1.00	42.21
	MOTA	3823	CG1	ILE	A	408	68.613	40.974	34.281	1.00	41.88
	MOTA	3824	CD1	ILE	A	408	67.762	39.937	34.976	1.00	46.39
50	MOTA	3825	С	ILE	A	408	71.480	40.861	31.746	1.00	39.97
	ATOM	3826	0	ILE	A	408	72.639	41.158	32.006	1.00	40.89
55	MOTA	3827	N	VAL	A	409	71.155	40.027	30.767	1.00	39.94
	MOTA	3829	CA	VAL	A	409	72.178	39.342	29.988	1.00	39.04
	MOTA	3830	СВ	VAL	A	409	71.560	38.354	28.961	1.00	38.16
60	MOTA	3831	CG1	VAL	A	409	72.656	37.687	28.142	1.00	37.45
	MOTA	3832	CG2	LAV	A	409	70.741	37.291	29.686	1.00	36.14
65	ATOM	3833	С	VAL	A	409	73.252	40.227	29.343	1.00	38.82
05	MOTA	3834	0	VAL	A	409	74.426	40.093	29.675	1.00	39.64

	ATOM	3835	N	PRO A 410	72.881	41.145	28.432	1.00	38.89
	MOTA	3836	CD	PRO A 410	71.583	41.362	27.770	1.00	38.55
5	MOTA	3837	CA	PRO A 410	73.917	41.984	27.818	1.00	38.75
	ATOM	3838	CB	PRO A 410	73.207	42.562	26.597	1.00	38.19
10	ATOM	3839	CG	PRO A 410	71.806	42.667	27.051	1.00	38.80
10	MOTA	3840	С	PRO A 410	74.535	43.083	28.685	1.00	40.43
	ATOM	3841	0	PRO A 410	75.607	43.593	28.359	1.00	41.27
15	ATOM	3842	N	ILE A 411	73.870	43.461	29.771	1.00	41.44
`.	ATOM	3844	CA	ILE A 411	74.404	44.503	30.649	1.00	41.02
	MOTA	3845	СВ	ILE A 411	73.285	45.380	31.239	1.00	40.99
20	MOTA	3846	CG2	ILE A 411	73.890	46.497	32.086	1.00	41.83
	ATOM	3847	CG1	ILE A 411	72.418	45.967	30.122	1.00	42.47
25	ATOM	3848	CD1	ILE A 411	73.112	47.005	29.264	1.00	45.28
	ATOM	3849	С	ILE A 411	75.212	43.923	31.809	1.00	40.15
20	MOTA	3850	0	ILE A 411	76.254	44.460	32.186	1.00	39.92
30	MOTA	3851	N	LEU A 412	74.716	42.831	32.379	1.00	39.13
	MOTA	3853	CA	LEU A 412	75.373	42.185	33.506	1.00	39.13
35	MOTA	3854	СВ	LEU A 412	74.335	41.817	34.570	1.00	39.71
	MOTA	3855	CG	LEU A 412	73.493	42.941	35.179	1.00	38.70
40	ATOM	3856	CD1	LEU A 412	72.548	42.362	36.220	1.00	36.24
40	MOTA	3857	CD2	LEU A 412	74.403	43.986	35.802	1.00	39.15
	MOTA	3858	С	LEU A 412	76.209	40.945	33.175	1.00	39.67
45	MOTA	3859	0	LEU A 412	77.322	40.796	33.677	1.00	41.17
	MOTA	3860	N	VAL A 413	75.678	40.060	32.336	1.00	38.13
50	ATOM	3862	CA	VAL A 413	76.369	38.818	31.991	1.00	37.18
50	MOTA	3863	СВ	VAL A 413	75.344	37.708	31.642	1.00	38.08
	MOTA	3864	CG1	VAL A 413	76.042	36.373	31.445	1.00	39.08
55	MOTA	3865	CG2	VAL A 413	74.303	37.591	32.753	1.00	35.49
	ATOM	3866	С	VAL A 413	77.477	38.893	30.926	1.00	36.23
60	MOTA	3867	0	VAL A 413	78.634	38.583	31.221	1.00	36.97
00	MOTA	3868	N	LEU A 414	77.134	39.296	29.702	1.00	34.70
	MOTA	3870	CA	LEU A 414	78.108	39.394	28.610	1.00	32.37
65	ATOM	3871	CB	LEU A 414	77.500	40.081	27.390	1.00	31.45
	ATOM	3872	CG	LEU A 414	77.451	39.326	26.062	1.00	31.86

	ATOM	3873	CD1	LEU A	414	77.167	40.338	24.952	1.00	34.05
5	MOTA	3874	CD2	LEU A	414	78.751	38.585	25.790	1.00	30.36
3	MOTA	3875	С	LEU A	414	79.403	40.109	28.972	1.00	33.92
	MOTA	3876	0	LEU A	414	80.485	39.643	28.619	1.00	36.82
10	MOTA	3877	N	PRO A	415	79.314	41.285	29.619	1.00	34.56
	ATOM	3878	CD	PRO A	415	78.103	42.090	29.858	1.00	34.83
15	ATOM	3879	CA	PRO A	415	80.503	42.046	30.011	1.00	35.76
15	MOTA	3880	CB	PRO A	415	79.915	43.149	30.883	1.00	35.11
	ATOM	3881	CG	PRO A	415	78.673	43.467	30.151	1.00	35.76
20	ATOM	3882	С	PRO A	415	81.583	41.263	30.748	1.00	36.22
	MOTA	3883	0	PRO A	415	82.760	41.387	30.418	1.00	37.32
25	MOTA	3884	N	ARG A	416	81.194	40.466	31.739	1.00	39.00
42	MOTA	3886	CA	ARG A	416	82.167	39.686	32.501	1.00	41.92
	ATOM	3887	CB	ARG A	416	81.517	39.047	33.731	1.00	46.13
30	ATOM	3888	CG	ARG A	416	81.111	40.038	34.823	1.00	52.86
	ATOM	3889	CD	ARG A	416	82.299	40.830	35.366	1.00	60.47
35	ATOM	3890	NE	ARG A	4 416	83.287	39.986	36.040	1.00	69.03
33	MOTA	3892	CŹ	ARG A	4 416	83.205	39.589	37.309	1.00	73.10
	ATOM	3893	NH1	ARG A	A 416	82.174	39.955	38.062	1.00	76.24
40	MOTA	3896	NH2	ARG A	A 416	84.163	38.833	37.834	1.00	73.45
	ATOM	3899	С	ARG Z	A 416	82.826	38.622	31.636	1.00	42.34
45	ATOM	3900	0	ARG I	A 416	84.031	38.405	31.717	1.00	43.31
•-	MOTA	3901	N	VAL 2	A 417	82.040	37.990	30.774		42.15
	MOTA	3903	CA	VAL	A 417	82.554	36.955	29.886	1.00	41.06
50	ATOM	3904	CB	VAL .	A 417	81.418	36.302	29.091	1.00	43.59
	ATOM	3905	CG1	. VAL .	A 417	81.976	35.261	28.127	1.00	48.62
55	ATOM	3906	CG2	VAL .	A 417	80.414	35.672	30.038	1.00	45.02
	ATOM	3907	С	VAL	A 417	83.581	37.500	28.902	1.00	38.49
	ATOM	3908	0	VAL	A 417	84.658	36.929	28.729	1.00	38.46
60	ATOM	3909	N	ASN	A 418	83.236	38.598	28.243	1.00	36.89
	ATOM	3911	CA	ASN	A 418	84.130	39.206	27.273	1.00	34.67
65	ATOM	3912	CB	asn	A 418	83.448	40.370	26.564	1.00	36.25
	ATOM	3913	CG	ASN	A 418	82.506	39.913	25.475	1.00	37.58

	MOTA	3914	OD1	ASN .	A	418	82.585	38.776	25.014	1.00	39.99-
	ATOM	3915	ND2	ASN .	A	418	81.613	40.796	25.055	1.00	36.92
5	MOTA	3918	С	ASN .	A	418	85.426	39.671	27.896	1.00	34.68
	MOTA	3919	0	ASN .	A	418	86.421	39.831	27.199	1.00	35.38
10	MOTA	3920	N	GLU .	A	419	85.426	39.895	29.206	1.00	35.06
10	ATOM	3922	CA	GLU	A	419	86.640	40.336	29.879	1.00	37.31
	ATOM	3923	CB	GLU	A	419	86.351	40.788	31.304	1.00	41.20
15	MOTA	3924	CG	GLU	A	419	87.557	41.433	31.972	1.00	49.99
	MOTA	3925	CD	GLU	A	419	87.312	41.800	33.419	1.00	54.63
20	MOTA	3926	OE1	GLU	Α	419	86.182	42.226	33.751	1.00	59.06
20	ATOM	3927	OE2	GLU	A	419	88.257	41.666	34.227	1.00	59.31
	MOTA	3928	С	GLU	A	419	87.682	39.222	29.889	1.00	37.58
25	MOTA	3929	0 (GLU	A	419	88.882	39.486	29.785	1.00	38.06
	ATOM	3930	N	LYS	Α	420	87.232	37.980	30.049	1.00	37.28
30	ATOM	3932	CA	LYS	A	420	88.154	36.853	30.033	1.00	37.63
30	ATOM	3933	CB	LYS	A	420	87.492	35.591	30.587	1.00	37.80
	MOTA	3934	CG	LYS	A	420	88.379	34.342	30.510	1.00	43.22
35	MOTA	3935	CD	LYS	A	420	89.698	34.521	31.265	1.00	46.31
	MOTA	3936	CE	LYS	A	420	90.650	33.352	31.031	1.00	48.33
40	ATOM	3937	NZ	LYS	A	420	91.993	33.578	31.644	1.00	48.07
40	MOTA	3941	С	LYS	A	420	88.610	36.631	28.593	1.00	38.72
	MOTA	3942	0	LYS	A	420	89.802	36.479	28.329	1.00	41.62
45	MOTA	3943	N	LEU	A	421	87.663	36.659	27.657	1.00	38.62
	MOTA	3945	CA	LEU	A	421	87.975	36.466	26.246	1.00	38.52
50	ATOM	3946	CB	LEU	A	421	86.705	36.474	25.395	1.00	36.49
	MOTA	3947	CG	LEU	A	421	85.770	35.278	25.539	1.00	34.05
	ATOM	3948	CD1	LEU	A	421	84.533	35.489	24.703	1.00	31.56
55	MOTA	3949	CD2	LEU	A	421	86.492	34.014	25.113	1.00	36.95
	MOTA	3950	С	ĻĚU	A	421	88.941	37.525	25.744	1.00	40.94
60	MOTA	3951	0	LEU	A	421	89.797	37.237	24.917	1.00	43.18
00	MOTA	3952	N	GLN	A	422	88.783	38.756	26.218	1.00	44.55
	MOTA	3954	CA	GLN	A	422	89.672	39.844	25.820	1.00	48.56
65	MOTA	3955	CB	GLN	A	422	89.036	41.210	26.101	1.00	52.32
	ATOM	3956	CG	GLN	A	422	88.278	41.821	24.914	1.00	58.07

	MOTA	3957	CD	GLN .	A	422	89.197	42.459	23.863	1.00	62.49
_	ATOM	3958	OE1	GLN 2	A	422	90.361	42.770	24.133	1.00	63.36
5	ATOM	3959	NE2	GLN :	A	422	88.661	42.673	22.665	1.00	63.04
	ATOM	3962	С	GLN .	A	422	91.037	39.725	26.501	1.00	49.05
10	ATOM	3963	0	GLN .	A	422	92.013	40.323	26.045	1.00	50.57
	MOTA	3964	N	LYS .	A	423	91.096	38.996	27.615	1.00	49.71
1.6	ATOM	3966	CA	LYS .	A	423	92.366	38.777	28.305	1.00	50.46
15	MOTA	3967	СВ	LYS .	A	423	92.151	38.125	29.670	1.00	52.85
	MOTA	3968	CG	LYS .	A	423	93.442	37.862	30.434	1.00	56.39
20	MOTA	3969	CD	LYS	A	423	93.156	37.388	31.845	1.00	59.26
	MOTA	3970	CE	LYS	A	423	92.354	38.432	32.611	1.00	60.80
26	MOTA	3971	NZ	LYS	A	423	92.070	37.998	34.006	1.00	64.66
25	ATOM	3975	С	LYS	A	423	93.108	37.827	27.378	1.00	49.36
	ATOM	3976	0	LYS	A	423	94.320	37.945	27.174	1.00	51.82
30	ATOM	3977	N	GLY	A	424	92.349	36.884	26.824	1.00	46.45
	ATOM	3979	CA	GLY	A	424	92.883	35.939	25.868	1.00	40.64
35	ATOM	3980	С	GLY	A	424	93.484	34.658	26.377	1.00	38.23
33	ATOM	3981	0	GLY	A	424	93.364	34.303	27.552	1.00	36.91
	MOTA	3982	N	PHE	A	425	94.111	33.950	25.445	1.00	36.95
40	MOTA	3984	CA	PHE	A	425	94.768	32.684	25.712	1.00	34.69
	MOTA	3985	CB	PHE	A	425	93.986	31.530	25.075	1.00	38.25
45	ATOM	3986	CG	PHE	A	425	92.570	31.402	25.571	1.00	44.42
	MOTA	3987	CD1	PHE	A	425	92.294	31.343	26.937	1.00	44.31
	ATOM	3988	CD2	PHE	A	425	91.508	31.345	24.671	1.00	46.34
50	ATOM	3989	CE1	PHE	A	425	90.986	31.235	27.399	1.00	45.62
	ATOM	3990	CE2	PHE	A	425	90.192	31.236	25.125	1.00	48.05
55	ATOM	3991	CZ	PHE	A	425	89.933	31.181	26.492	1.00	46.50
23	ATOM	3992	С	PHE	A	425	96.153	32.748	25.091	1.00	31.44
	MOTA	3993	0	PHE	A	425	96.316	33.236	23.982	1.00	31.46
60	ATOM	3994	N	PRO	A	426	97.176	32.316	25.831	1.00	29.84
	MOTA	3995	CD	PRO	A	426	97.076	31.978	27.259	1.00	30.24
65	MOTA	3996	CA	PRO	A	426	98.571	32.299	25.396	1.00	.30.20
U.J	ATOM	3997	СВ	PRO	A	426	99.269	31.630	26.570	1.00	28.60

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	MOTA	3998	CG	PRO A 426	98.494	32.133	27.725	1.00 31.57
	MOTA	3999	С	PRO A 426	98Ļ777	31.489	24.123	1.00 31.10
5	MOTA	4000	0	PRO A 426	98.258	30.377	23.998	1.00 34.08
	ATOM	4001	N	LEU A 427	99.570	32.033	23.203	1.00 29.97
10	MOTA	4003	CA	LEU A 427	99.878	31.377	21.935	1.00 29.50
10	ATOM	4004	СВ	LEU A 427	99.991	32.433	20.831	1.00 28.79
	MOTA	4005	CG	LEU A 427	99.106	32.365	19.576	1.00 31.67
15	ATOM	4006	CD1	LEU A 427	97.713	31.822	19.854	1.00 29.49
	MOTA	4007	CD2	LEU A 427	99.020	33.753	18.977	1.00 30.38
20	MOTA	4008	С	LEU A 427	101.189	30.602	22.095	1.00 28.95
20	ATOM	4009	0	LEU A 427	102.012	30.939	22.943	1.00 30.17
	MOTA	4010	N	PRO A 428	101.396	29.548	21.293	1.00 29.54
25	MOTA	4011	CD	PRO A 428	100.502	29.048	20.234	1.00 29.81
	MOTA	4012	CA	PRO A 428	102.614	28.734	21.368	1.00 31.77
30	MOTA	4013	СВ	PRO A 428	102.218	27.492	20.584	1.00 30.20
30	MOTA	4014	CG	PRO A 428	101.382	28.073	19.490	1.00 27.41
	MOTA	4015	С	PRO A 428	103.843	29.405	20.753	1.00 35.74
35	MOTA	4016	0	PRO A 428	104.361	28.950	19.731	1.00 39.47
	MOTA	4017	N	THR A 429	104.323	30.472	21.379	1.00 36.82
40	MOTA	4019	CA	THR A 429	105.486	31.189	20.866	1.00 36.58
	ATOM	4020	CB	THR A 429	105.200	32.694	20.735	1.00 -38.98
	MOTA	4021	OG1	THR A 429	104.616	33.183	21.953	1.00 37.38
45	MOTA	4023	CG2	THR A 429	104.262	32.956	19.565	1.00 41.95
•	ATOM	4024	С	THR A 429	106.686	31.024	21.769	1.00 35.72
50	MOTA	4025	0	THR A 429	106.567	31.108	22.994	1.00 35.83
	MOTA	4026	N	PRO A 430	107.867	30.791	21.182	1.00 34.72
	MOTA	4027	CD	PRO A 430	108.180	30.643	19.753	1.00 36.21
55	MOTA	4028	CA	PRO A 430	109.069	30.629	21.995	1.00 34.61
	MOTA	4029	CB	PRO A 430	110.124		20.963	1.00 33.90
60	MOTA	4030	CG	PRO A 430	109.658	30.907	19.726	1.00 33.59
	MOTA	4031	С	PRO A 430	109.388		22.691	1.00 35.01
	MOTA	4032	0	PRO A 430	108.648			
65	MOTA	4033	N	ALA A 431	110.457			
	MOTA	4035	CA	ALA A 431	110.824	33.190	24.173	1.00 36.53

	MOTA	4036	CB	ALA A	A	431	111.801	32.878	25.298	1.00	36.36
5	MOTA	4037	С	ALA 2	A	431	111.416	34.241	23.243	1.00	38.15
3	ATOM	4038	0	ALA :	A	431	111.944	33.932	22.174	1.00	39.81
	MOTA	4039	N	ARG .	A	432	111.270	35.495	23.641	1.00	40.43
10	MOTA	4041	CA	ARG .	A	432	111.819	36.620	22.902	1.00	41.66
	MOTA	4042	СВ	ARG .	A	432	113.349	36.537	22.887	1.00	44.65
15	ATOM	4043	CG	ARG .	A	432	113.990	36.686	24.262	1.00	51.96
15	MOTA	4044	CD	ARG .	A	432	115.516	36.654	24.185	1.00	59.79
	ATOM	4045	NE	ARG .	A	432	116.029	35.372	23.700	1.00	63.72
20	ATOM	4047	CZ	ARG .	A	432	117.302	34.993	23.777	1.00	67.96
·	ATOM	4048	NH1	ARG	A	432	118.212	35.795	24.320	1.00	70.88
25	ATOM	4051	NH2	ARG	A	432	117.668	33.804	23.314	1.00	68.07
25	ATOM	4054	С	ARG	A	432	111.293	36.898	21.499	1.00	40.87
	ATOM	4055	0	ARG	A	432	111.963	37.567	20.721	1.00	42.99
30	ATOM	4056	N	VAL	A	433	110.103	36.414	21.166	1.00	40.50
	MOTA	4058	CA	VAL	A	433	109.538	36.693	19.847	1.00	39.69
35	ATOM	4059	CB	VAL	A	433	109.325	35.415	18.982	1.00	39.93
<i>JJ</i>	ATOM	4060	CG1	VAL	A	433	110.588	34.575	18.943	1.00	39.03
	ATOM	4061	CG2	VAL	A	433	108.134	34.608	19.470	1.00	39.91
40	MOTA	4062	С	VAL	A	433	108.206	37.399	20.024	1.00	41.27
	MOTA	4063	0	VAL	A	433	107.538	37.240	21.048	1.00	43.57
45	MOTA	4064	N	GLN	A	434	107.845	38.223	19.053	1.00	41.32
	MOTA	4066	CA	GLN	A	434	106.583	38.939	19.104	1.00	41.91
	ATOM	4067	CB	GLN	A	434	106.802	40.411	19.454	1.00	45.76
50	ATOM	4068	CG	GLN	A	434	105.706	41.018	20.335	1.00	52.61
	ATOM	4069	CD	GLN	A	434	105.850	40.687	21.829	1.00	56.74
55	ATOM	4070	OE1	GLN	A	434	105.372	41.436	22.681	1.00	58.13
	ATOM	4071	NE2	GLN	A	434	106.512	39.579	22.149	1.00	57.45
	MOTA	4074	С	GLN	A	434	106.005	38.804	17.716	1.00	40.30
60	ATOM	4075	0	GLN	A	434	106.745	38.821	16.732	1.00	40.86
	ATOM	4076	N	LEU	A	435	104.695	38.614	17.642	1.00	39.20
65	MOTA	4078	CA	LEU	A	435	104.018	38.439	16.366		39.99
	ATOM	4079	CB	LEU	A	435	102.965	37.338	16.501	1.00	38.11

	ATOM	4080	CG	LEU A	435	103.347	36.088	17.309	1.00	37.55-
	ATOM	4081	CD1	LEU A	435	102.118	35.238	17.538	1.00	37.75
5	ATOM	4082	CD2	LEU A	435	104.426	35.281	16.613	1.00	36.46
	ATOM	4083	С	LEU A	435	103.358	39.739	15.916	1.00	41.18
10	ATOM	4084	0	LEU A	435	102.660	40.387	16.695	1.00	42.90
10	ATOM	4085	N	TYR A	436	103.590	40.125	14.665	1.00	42.57
	MOTA	4087	CA	TYR A	436	103.004	41.348	14.116	1.00	43.80
15	MOTA	4088	СВ	TYR A	436	104.021	42.510	14.120	1.00	43.48
	ATOM	4089	CG	TYR A	436	105.232	42.328	13.224	1.00	43.66
20	ATOM	4090	CD1	TYR A	436	106.259	41.452	13.573	1.00	43.96
20	ATOM	4091	CE1	TYR A	436	107.364	41.271	12.743	1.00	44.63
	MOTA	4092	CD2	TYR A	436	105.345	43.028	12.020	1.00	43.04
25	MOTA	4093	CE2	TYR A	436	106.448	42.852	11.181	1.00	43.53
	MOTA	4094	cz	TYR A	436	107.452	41.970	11.549	1.00	44.29
30	MOTA	4095	ОН	TYR A	436	108.528	41.757	10.711	1.00	44.66
30	MOTA	4097	С	TYR A	436	102.457	41.090	12.712	1.00	43.77
	MOTA	4098	0	TYR A	436	102.729	40.046	12.122	1.00	45.49
35	MOTA	4099	N	ASN A	437	101.687	42.042	12.189	1.00	45.09
	MOTA	4101	CA	ASN A	437	101.069	41.932	10.865	1.00	45.97
40	MOTA	4102	CB	ASN A	437	102.083	42.231	9.750	1.00	51.76
,,,	MOTA	4103	CG	ASN A	437	102.161	43.720	9.416	1.00	57.75
	MOTA	4104	OD1	ASN A	437	102.832	44.492	10.105	1.00	59.59
45	MOTA	4105	ND2	ASN A	437	101.450	44.131	8.370	1.00	60.25
	ATOM	4108	C	ASN A	437	100.385	40.581	10.665	1.00	44.47
50	MOTA	4109	0	ASN A	437	100.743	39.797	9.786	1.00	44.02
	MOTA	4110	N.	VAL A	438	99.360	40.344	11.475	1.00	44.00
	MOTA	4112	CA	VAL A	438	98.622	39.091	11.444	1.00	43.09
55	MOTA	4113	CB	VAL A	438	98.214	38.646	12.871	1.00	43.35
	MOTA	4114	CG1	VAL A	438	99.425	38.629	13.786	1.00	43.74
60	MOTA	4115	CG2	VAL A	438	97.143	39.562	13.433	1.00	43.99
	MOTA	4116	С	VAL A	438	97.374	39.101	10.579	1.00	41.42
	ATOM	4117	0	VAL A	438	96.740	40.139	10.377	1.00	39.58
65	ATOM	4118	N	VAL A	439	97.018	37.912	10.110	1.00	41.86
	MOTA	4120	CA	VAL A	439	95.839	37.689	9.291	1.00	43.18

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7.829 1.00 42.54 MOTA 4121 CB VAL A 439 96.202 37.377 96.568 38.646 MOTA 4122 CG1 VAL A 439 7.104 1.00 46.06 5 MOTA 4123 CG2 VAL A 439 97.351 36.391 7.770 1.00 41.82 С **VAL A 439** 95.117 36.482 9.866 1.00 43.37 ATOM 4124 10 **ATOM** 0 **VAL A 439** 95.733 35.446 10.103 1.00 45.02 4125 **LEU A 440** 93.829 36.631 10.141 1.00 43.45 **ATOM** 4126 N **ATOM** 4128 CA **LEU A 440** 93.041 35.535 10.681 1.00 42.92 15 CB **LEU A 440** 92.355 35.956 11.978 1.00 42.49 MOTA 4129 4130 CG **LEU A 440** 93.220 36.062 13.234 1.00 42.93 MOTA 20 CD1 LEU A 440 93.920 1.00 44.28 MOTA 4131 34.745 13.462 CD2 LEU A 440 94.231 37.181 13.111 1.00 45.59 ATOM 4132 MOTA 4133 C LEU A 440 92.008 35.132 9.643 1.00 44.03 25 **ATOM** 4134 0 **LEU A 440** 91.017 35.831 9.446 1.00 45.13 **GLN A 441** 92.268 34.029 8.947 1.00 44.00 ATOM 4135 N 30 4137 **GLN A 441** 91.363 1.00 43.66 ATOM CA 33.536 7.910 **GLN A 441** 92.148 6.678 1.00 48.64 **ATOM** 4138 CB 33.058 MOTA 4139 CG **GLN A 441** 92.881 34.113 5.860 1.00 54.68 35 4.627 93.556 33.515 1.00 57.71 **ATOM** 4140 CD GLN A 441 OE1 GLN A 441 93.111 4.093 1.00 59.57 **ATOM** 4141 32.495 40 **ATOM** 4142 NE2 GLN A 441 94.637 34.141 4.179 1.00 60.18 MOTA 4145 C **GLN A 441** 90.528 32.355 8.390 1.00 41.14 91.055 31.249 8.541 1.00 41.88 ATOM 4146 0 **GLN A 441** 45 1.00 38.75 MOTA 4147 N PRO A 442 89.217 32.553 8.613 MOTA 4148 CD PRO A 442 88.444 33.805 8.553 1.00 39.00 50 CA PRO A 442 88.372 31.442 9.065 1.00 35.99 MOTA 4149 ATOM 4150 CB PRO A 442 87.049 32.131 9.381 1.00 36.27 ATOM 4151 CG PRO A 442 87.038 33.295 8.450 1.00 38.58 55 ATOM 4152 C PRO A 442 88.234 30.403 7.952 1.00 34.89 ATOM 4153 0 PRO A 442 88.109 30.751 6.780 1.00 37.34 60 **ATOM** 4154 N HIS A 443 88.332 29.130 8.317 1.00 32.88 MOTA CA HIS A 443 88.242 28.036 7.361 1.00 30.71 4156 89.610 ATOM 4157 CB HIS A 443 27.377 7.172 1.00 31.17 65 **ATOM** 4158 CG HIS A 443 90.508 28.103 6.225 1.00 29.53

	MOTA	4159	CD2	HIS A	443	91.098	27.702	5.076	1.00	27.37-
	MOTA	4160	ND1	HIS A	443	90.868	29.421	6.400	1.00	31.12
5	MOTA	4162	CEI	HIS A	443	91.635	29.804	5.397	1.00	30.58
	MOTA	4163	NE2	HIS A	443	91.790	28.779	4.580	1.00	29.90
10	MOTA	4165	С	HIS A	443	87.263	27.003	7.863	1.00	31.31
10	MOTA	4166	0	HIS A	443	86.698	27.157	8.936	1.00	36.03
•	MOTA	4167	N	GLN A	444	87.058	25.948	7.086	1.00	33.00
15	ATOM	4169	CA	GLN A	444	86.136	24.896	7.481	1.00	33.08
	MOTA	4170	CB	GLN A	444	85.802	24.000	6.282	1.00	35.23
20	ATOM	4171	CG	GLN A	444	84.829	22.860	6.570	1.00	37.52
20	MOTA	4172	CD	GLN A	444	83.436	23.337	6.954	1.00	41.28
	ATOM	4173	OE1	GLN A	444	83.142	24.531	6.941	1.00	42.98
25	MOTA	4174	NE2	GLN A	444	82.570	22.395	7.298	1.00	44.68
	ATOM	4177	С	GLN A	444	86.735	24.069	8.613	1.00	32.22
30	ATOM	4178	0	GLN A	444	87.720	23.360	8.416	1.00	34.24
30	ATOM	4179	N	asn a	445	86.175	24.230	9.808	1.00	29.86
	ATOM	4181	CA	ASN A	445	86.584	23.492	11.003	1.00	28.34
35	ATOM	4182	CB	asn a	445	86.628	21.984	10.714	1.00	27.65
	MOTA	4183	CG	ASN A	445	85.272	21.427	10.319	1.00	28.84
40	ATOM	4184	OD1	ASN A	445	85.155	20.637	9.385	1.00	33.08
40	MOTA	4185	ND2	ASN A	445	84.233	21.850	11.022	1.00	32.64
	MOTA	4188	С	ASN A	445	87.863	23.950	11.709	1.00	27.86
45	MOTA	4189	0	ASN A	445	88.318	23.306	12.658	1.00	30.31
	ATOM	4190	N	PHE A	446	88.436	25.064	11.272	1.00	26.55
50	MOTA	4192	CA	PHE A	446	89.639	25.582	11.904	1.00	25.16
	ATOM	4193	CB	PHE A	446	90.858	24.688	11.626	1.00	28.26
	MOTA	4194	CG	PHE A	446	91.407	24.796	10.233	1.00	29.75
55	MOTA	4195	CD1	PHE A	446	90.953	23.956	9.223	1.00	29.14
	MOTA	4196	CD2	PHE A	446	92.413	25.711	9.940	1.00	32.05
60	ATOM	4197	CE1	PHE A	446	91.492	24.020	7.950	1.00	26.24
00	ATOM	4198	CE2	PHE A	446	92.961	25.784	8.662	1.00	30.94
	MOTA	4199	CZ	PHE A	446	92.500	24.936	7.667	1.00	29.55
65	MOTA	4200	С	PHE A	. 446	89.906	27.020	11.506	1.00	25.07
	MOTA	4201	0	PHE A	446	89.372	27.508	10.513	1.00	24.16

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	ATOM	4202	N	LEU A	447	90.718	27.697	12.306	1.00 26.36
_	MOTA	4204	CA	LEU A	A 447	91.059	29.091	12.076	1.00 25.79
5	ATOM	4205	СВ	LEU A	A 447	90.825	29.880	13.365	1.00 27.64
	MOTA	4206	CG	LEU A	A 447	91.097	31.384	13.422	1.00 28.75
10	MOTA	4207	CD1	LEU A	A 447	90.193	32.125	12.449	1.00 27.40
	ATOM	4208	CD2	LEU A	A 447	90.863	31.875	14.843	1.00 25.78
15	MOTA	4209	С	LEU A	A 447	92.508	29.221	11.635	1.00 25.68
15	ATOM	4210	0	LEU A	A 447	93.398	28.619	12.223	1.00 26.90
	ATOM	4211	N	LEU A	A 448	92.738	30.000	10.589	1.00 26.87
20	MOTA	4213	CA	LEU A	A 448	94.080	30.199	10.084	1.00 26.96
	MOTA	4214	СВ	LEU A	A 448	94.074	30.247	8.559	1.00 26.42
25	MOTA	4215	CG	LEU A	A 448	95.432	30.045	7.886	1.00 25.05
25	MOTA	4216	CD1	LEU A	A 448	96.040	28.738	8.331	1.00 25.88
	MOTA	4217	CD2	LEU 1	A 448	95.273	30.049	6.396	1.00 29.10
30	MOTA	4218	С	LEU A	A 448	94.627	31.493	10.651	1.00 29.88
	MOTA	4219	0	LEU A	A 448	94.005	32.552	10.530	1.00 28.70
35	MOTA	4220	N	PHE A	A 449	95.788	31.386	11.283	1.00 32.23
))	ATOM	4222	CA	PHE 2	A 449	96.469	32.511	11.897	1.00 32.39
	ATOM	4223	СВ	PHE 2	A 449	96.655	32.234	13.394	1.00 31.41
40	MOTA	4224	CG	PHB 2	A 449	97.511	33.243	14.100	1.00 31.44
	ATOM	4225	CD1	PHE I	A 449	98.897	33.160	14.046	1.00 32.18
45	ATOM	4226	CD2	PHE 2	A 449	96.934	34.296	14.797	1.00 35.10
45	ATOM	4227	CE1	PHE 2	A 449	99.695	34.111	14.671	1.00 32.81
	ATOM	4228	CE2	PHE 2	A 449	97.725	35.257	15.429	1.00 34.26
50	ATOM	4229	CZ	PHE .	A 449	99.106	35.162	15.363	1.00 33.12
	MOTA	4230	С	PHE :	A 449	97.829	32.685	11.228	1.00 34.46
. 55	MOTA	4231	0	PHE .	A 449	98.717	31.865	11.420	1.00 38.92
	MOTA	4232	N	GLY .	A 450	97.990	33.738	10.439	1.00 34.70
	MOTA	4234	CA	GLY .	A 450	99.262	33.982	9.781	1.00 33.42
60	MOTA	4235	С	GLY .	A 450	99.901	35.207	10.397	1.00 35.08
	MOTA	4236	0	GLY .	A 450	99.194	36.140	10.784	1.00 36.42
65	ATOM	4237	N	ALA .	A 451	101.225	35.234	10.491	1.00 33.89
JJ	MOTA	4239	CA	ALA	A 451	101.889	36.383	11.092	1.00 33.95

	MOTA	4240	СВ	ALA	A	451	101.649	36.384	12.587	1.00	31.78
	ATOM	4241	С	ALA	A	451	103.382	36.454	10.815	1.00	34.43
5	ATOM	4242	0	ALA	Α	451	103.986	35.489	10.335	1.00	36.02
	ATOM	4243	N	ASP	A	452	103.954	37.628	11.055	1.00	33.44
10	ATOM	4245	CA	ASP	Α	452	105.381	37.847	10.885	1.00	35.01
10	ATOM	4246	CB	ASP	A	452	105.670	39.186	10.196	1.00	37.76
	MOTA	4247	CG	ASP	A	452	105.437	39.144	8.696	1.00	40.65
15	MOTA	4248	OD1	ASP	A	452	105.743	38.106	8.074	1.00	42.78
	ATOM	4249	OD2	ASP	A	452	104.961	40.159	8.134	1.00	42.90
20	MOTA	4250	С	ASP	A	452	105.913	37.875	12.306	1.00	35.28
20	MOTA	4251	0	ASP	Α	452	105.196	38.247	13.237	1.00	33.85
	MOTA	4252	N	VAL	A	453	107.174	37.509	12.473	1.00	37.44
25	MOTA	4254	CA	VAL	A	453	107.771	37.468	13.798	1.00	38.73
	MOTA	4255	CB	VAL	A	453	108.253	36.034	14.138	1.00	39.37
30	MOTA	4256	CG1	VAL	A	453	108.678	35.955	15.591	1.00	40.45
50	MOTA	4257	CG2	VAL	A	453	107.169	35.004	13.819	1.00	37.45
	MOTA	4258	С	VAL	A	453	108.970	38.398	13.899	1.00	39.49
35	MOTA	4259	0	VAL	A	453	109.669	38.641	12.914	1.00	40.46
	MOTA	4260	N	VAL	A	454	109.190	38.929	15.092	1.00	39.41
40	MOTA	4262	CA	VAL	A	454	110.324	39.796	15.331	1.00	40.55
	MOTA	4263	СВ	VAL	A	454 ´	109.895	41.280	15.511	1.00	41.60
	MOTA	4264	CG1	VAL	A	454	108.859	41.423	16.605	1.00	42.90
45	MOTA	4265	CG2	VAL	A	454	111.105	42.149	15.798	1.00	43.42
	MOTA	4266	С	VAL	A	454	111.037	39.256	16.561		41.20
50	ATOM	4267	0	VAL	A	454	110.440	39.115	17.626		41.61
	ATOM	4268	N			455	112.284	38.851	16.376		42.91
	ATOM	4270	CA			455	113.084	38.309	17.460		45.62
55	ATOM	4271	CB			455	114.015	37.227	16.908		44.83
	ATOM	4272	CG			455	115.038	36.684	17.881		44.58
60	ATOM	4273		TYR			114.682	35.774	18.875		43.02
	ATOM	4274		TYR			115.644	35.246	19.739		44.27
	MOTA	4275		TYR			116.377	37.053	17.778		46.42
65	ATOM	4276		TYR			117.341	36.535	18.631		46.69
	MOTA	4277	CZ	TYR	A	455	116.973	35.634	19.607	1.00	46.43

	MOTA	4278	OH	TYR	A	455	117.944	35.141	20.445	1.00	49.56
5	MOTA	4280	С	TYR	A	455	113.864	39.443	18.128	1.00	49.88
3	MOTA	4281	ο .	TYR	A	455	114.558	40.213	17.455	1.00	49.95
•	MOTA	4282	N	LYS	A	456	113.694	39.552	19.444	1.00	54.41
10	MOTA	4284	CA	LYS	A	456	114.337	40.570	20.270	1.00	59.38
	MÕTA	4285	СВ	LYS	A	456	113.760	40.514	21.693	1.00	61.68
15	ATOM	4286	CG	LYS	A	456	114.401	41.475	22.692	1.00	66.56
15	MOTA	4287	CD	LYS	A	456	114.016	41.153	24.140	1.00	68.38
	ATOM	4288	CE	LYS	A	456	112.538	41.396	24.409	1.00	70.09
20	ATOM	4289	NZ	LYS	A	456	112.154	41.067	25.812	1.00	71.48
	MOTA	4293	С	LYS	A	456	115.855	40.382	20.311	1.00	62.37
25	MOTA	4294	OTI	LYS	A	456	116.324	39.430	20:976	1.00	64.57
23	ATOM	4295	OT2	LYS	A	456	116.559	41.196	19.673	1.00	66.00
	ATOM	4296	Cl	PC	A	777	121.817	32.468	19.343	1.00	72.53
30	MOTA	4297	C2	PC	A	777	121.094	31.121	19.465	1.00	63.82
	MOTA	4298	C3	PC	A	777	119.917	31.017	18.492	1.00	59.16
35	ATOM	4299	C4	PC	A	777	123.302	33.354	23.035	1.00	94.69
55	ATOM	4300	C5	PC	A	7 77	124.151	34.553	23.445	1.00	98.10
	MOTA	4301	C6	PC	A	777	125.683	33.126	24.652	1.00	99.68
40	ATOM	4302	C7	PC	A	777	126.361	35.328	24.003	1.00	100.00
	MOTA	4303	C8	PC	A	777	126.086	33.627	22.335	1.00	100.00
45	MOTA	4304	C31	PC	A	777	122.736	29.776	18.047	1.00	54.84
	MOTA	4305	C32	PC	A	777	122.337	30.654	16.869	1.00	51.64
	MOTA	4306	C33	PC	A	777	122.254	29.878	15.549	1.00	47.76
50	MOTA	4307	C34	PC	A	777	123.590	29.237	15.169	1.00	42.03
	MOTA	4308	C35	PC	A	777	123.429	28.173	14.085	1.00	39.41
55	MOTA	4309	C36	PC	A	777	122.582	28.674	12.916	1.00	36.18
	MOTA	4310	C37	PC	A	777	121.581	27.622	12.463	1.00	33.16
	MOTA	4311	C38	PC	A	777	120.679	28.158	11.377	1.00	31.40
60	MOTA	4312	C39	PC	A	777	119.610	29.062	11.951	1.00	36.47
	MOTA	4313	C40	PC	A	777	118.543	28.263	12.682	1.00	40.19
65	MOTA	4314	C41	PC	A	777	117.191	28.383	12.004	1.00	39.41
	MOTA	4315	C42	PC	A	777	116.468	27.050	11.956	1.00	41.86

	ATOM	4316	C43	PC	A	777	115.409	26.958	13.041	1.00	43.49-
	MOTA	4317	C44	PC	A	777	114.326	25 951	12.669	1.00	46.22
5	MOTA	4318	C45	PC	A	777	113.126	26.048	13.604	1.00	47.77
	MOTA	4319	C46	PC	A	777	112.271	24.786	13.561	1.00	48.67
10	ATOM	4320	C47	PC	A	777	111.948	24.277	14.964	1.00	48.62
10	ATOM	4321	C48	PC	A	777	110.492	24.449	15.368	1.00	45.72
	ATOM	4322	C11	PC	A	7 77	118.509	32.389	17.081	1.00	49.62
15	ATOM	4323	C12	PC	A	777 .	117.647	31.121	17.072	1.00	45.91
	ATOM	4324	· C13	PC	A	777	116.542	31.091	16.014	1.00	42.05
20	ATOM	4325	C14	PC	A	777	115.753	32.392	15.973	1.00	37.88
20	MOTA	4326	C15	PC	A	777	114.269	32.149	16.004	1.00	39.34
	ATOM	4327	C16	PC	A	777	113.573	32.877	14.874	1.00	39.38
25	ATOM	4328	C17	PC	A	7 7 7	112.066	32.688	14.951	1.00	43.09
	ATOM	4329	C18	PC	A	777	111.446	32.535	13.566	1.00	45.42
30	ATOM	4330	C19	PC	A	777	111.500	31.093	13.082	1.00	47.16
30	ATOM	4331	C20	PC	A	777	110.158	30.413	13.236	1.00	46.62
	MOTA	4332	C21	PC	A	777	109.917	30.004	14.673	1.00	46.65
35	ATOM	4333	C22	PC	A	777	108.444	29.836	14.942	1.00	46.97
	ATOM	4334	C23	PC	A	777	107.922	30.916	15.868	1.00	47.30
40	ATOM	4335	C24	PC	A	777	106.409	30.835	16.001	1.00	50.05
	ATOM	4336	C25	PC	A	777	105.996	29.898	17.121	1.00	50.59
	MOTA	4337	C26	PC	A	777	105.783	28.490	16.612	1.00	53.56
45	MOTA	4338	C27	PC	A	777	106.972	27.593	16.941	1.00	56.03
	MOTA	4339	C28	PC	A	777	107.348	26.625	15.831	1.00	55.75
50	MOTA	4340	011	PC	A	777	118.194	33.427	16.467	1.00	52.98
	MOTA	4341	031	PC	A	777	123.576	28.862	17.906	1.00	56.48
	MOTA	4342	02	PC	A	777	122.068	30.020	19.350	1.00	61.36
55	ATOM	4343	03	PC	A	777	119.731	32.307	17.866	1.00	53.04
	MOTA	4344	01P	PC	A	777	121.661	35.196	20.857	1.00	89.42
60	MOTA	4345	02P	PC	A	777	120.059	33.339	21.280	1.00	90.46
00	MOTA	4346	03P	PC	A	777	122.419	32.815	20.622	1.00	84.15
	MOTA	4347	04P	PC	A	777	121.916	33.769	22.889	1.00	91.71
65	MOTA	4348	N	PC	A	777	125.571	34.153	23.607	1.00	99.82
	ATOM	4349	P	PC	A	777	121.516	33.779	21.409	1.00	89.00

	ATOM	4350	Cl	PC	A	778	87.797	32.791	33.686	1.00 88.29
-	ATOM	4351	C2	PC	A	778	86.274	32.868	33.502	1.00 83.64
5	ATOM	4352	C3	PC	A	778	85.739	31.590	32.830	1.00 80.89
	ATOM	4353	C31	PC	A	778	84.282	33.619	34.787	1.00 76.37
10	MOTA	4354	C32	PC	A	778	83.704	33.859	33.391	1.00 70.52
	MOTA	4355	C33	PC	A	778	82.344	33.215	33.178	1.00 63.44
16	ATOM	4356	C34	PC	A	778	81.311	34.250	32.803	1.00 56.98
15	MOTA	4357	C35	PC	A	778	80.231	34.364	33.854	1.00 51.89
	MOTA	4358	C36	PC	A	778	79.088	33.400	33.593	1.00 48.39
20	ATOM	4359	C37	PC	A	778	78.611	33.453	32.155	1.00 47.51
	MOTA	4360	C38	PC	A	778	77.593	32.352	31.868	1.00 47.46
25	MOTA	4361	C39	PC	A	778	78.231	31.153	31.171	1.00 45.28
25	MOTA	4362	C40	PC	A	778	78.154	31.261	29.654	1.00 43.25
	MOTA	4363	C41	PC	A	778	79.381	31.957	29.099	1.00 40.88
30	ATOM	4364	C42	PC	A	778	79.324	32.082	27.589	1.00 42.02
	ATOM	4365	C43	PC	A	778	78.064	32.803	27.130	1.00 43.52
35	MOTA	4366	C44	PC	A	778	77.971	34.223	27.684	1.00 42.21
33	MOTA	4367	C45	PC	A	778	76.563	34.546	28.171	1.00 42.30
	MOTA	4368	C46	PC	A	778	75.523	34.335	27.077	1.00 42.97
40	MOTA	4369	C47	PC	A	778	74.121	34.185	27.662	1.00 42.80
	MOTA	4370	C48	PC	A	778	73.131	33.504	26.727	1.00 43.31
45	MOTA	4371	C11	. PC	A	. 778	86.101	31.542	30.412	1.00 71.27
.5	MOTA	4372	C12	PC	P	778	84.804	32.359	30.441	1.00 66.82
	ATOM	4373	C13	PC	P	778	83.700	31.845	29.548	1.00 59.12
50	MOTA	4374	C14	PC	I	778	83.836	32.380	28.141	1.00 55.92
	ATOM	4375	C15	5 PC	7	778	82.896	31.671	27.184	1.00 53.76
55	ATOM	4376	C16	5 PC	I	778	82.613	30.248	27.633	. 1.00 52.97
55	ATOM	4377	C1.	7 PC	I	778	83.562	29.250	26.990	1.00 50.51
	MOTA	4378	C18	B PC	1	A 778	84.990	29.415	27.480	1.00 47.60
60	MOTA	4379	C19	9 PC	7	778	85.860	30.046	26.408	1.00 47.69
	MOTA	4380	C20	0 · PC	. 1	778	86.359	29.011	25.411	1.00 44.90
65	MOTA	4381	C2:	1 PC	1	A 778	85.271	28.598	24.439	1.00 44.44
65	ATOM	4382	C2:	2 PC	7	A 778	85.574	29.064	23.038	1.00 44.15

	ATOM	4383	C23	PC	A	778	86.167	30.453	23.044	1.00 43.99
	ATOM	4384	C24	PC	A	778	87.670	30.411	22.853	1.00 42.63
5	ATOM	4385	C25	PC	A	778	88.068	31.161	21.599	1.00 43.28
	MOTA	4386	C26	PC	A	778	87.733	32.642	21.693	1.00 41.66
10	MOTA	4387	C27	PC	A	778	88.852	33.492	21.119	1.00 40.33
10	ATOM	4388	C28	PC	A	778	90.094	33.503	21.969	1.00 37.09
	MOTA	4389	011	PC	A	778	86.692	31.253	29.343	1.00 72.85
15	MOTA	4390	031	PC	A	778	83.619	33.857	35.833	1.00 78.26
	MOTA	4391	02	PC	A	778	85.652	33.065	34.823	1.00 81.22
20	ATOM	4392	03	PC	A	778	86.615	31.154	31.736	1.00 75.71
20	ATOM	4393	O1P	PC	A	778	89.401	31.823	36.960	1.00100.00
	MOTA	4394	O2P	PC	A	778	89.563	33.853	35.511	1.00100.00
25	MOTA	4395	03P	PC	A	778	88.125	31.917	34.809	1.00 96.24
	MOTA	4396	04 P	PC	A	778	87.424	33.342	36.757	1.00100.00
30	ATOM	4397	P	PC	A	778	88.628	32.739	36.008	1.00100.00
50	ATOM	4398	OH2	нон	A	901	116.815	15.747	16.799	1.00 43.90
	ATOM	4401	OH2	нон	A	902	107.439	22.033	22.778	1.00 44.99
35	ATOM	4404	OH2	нон	A	903	115.201	26.524	27.768	1.00 47.84
	MOTA	4407	OH2	нон	A	904	83.653	23.737	13.286	1.00 22.80
40	ATOM	4410	OH2	нон	A	905	76.576	23.779	22.886	1.00 55.25
	MOTA	4413	OH2	нон	A	906	101.110	43.595	19.882	1.00 44.52
	MOTA	4416	OH2	HOH	A	907	80.990	19.920	23.229	1.00 48.61
45	MOTA	4419	OH2	нон	A	908	141.374	35.733	18.039	1.00 56.87
	MOTA	4422	OH2	нон	A	909	81.958	22.677	23.953	1.00 61.10
50	MOTA	4425				910	152.059	36.697	-0.688	1.00 62.06
	MOTA	4428				911	139.649			
	MOTA	4431				912	124.594			
55	MOTA	4434				913	121.471			
	MOTA	4437					114.402		0.380	1.00 31.16
60	ATOM	4440				915				
	MOTA	4443					74.995			
<i>(</i>	ATOM	4446					105.633			
65	MOTA	4449				918	71.679			
	MOTA	4452	OH2	нон	A	919	104.762	14.189	10.052	1.00 57.21

	ATOM	4455	ОН2 НОН А 920	80.378	28.515	41.681	1.00 81.97
_	ATOM	4458	ОН2 НОН А 921	80.215	25.918	35.227	1.00 29.12
5	ATOM	4461	ОН2 НОН А 922	79.054	35.414	40.975	1.00 41.95
	ATOM	4464	ОН2 НОН А 923	65.692	27.690	51.372	1.00 65.22
10	ATOM	4467	ОН2 НОН А 924	88.914	24.630	4.689	1.00 48.96
	ATOM	4470	ОН2 НОН А 925	100.082	22.695	34.405	1.00 48.58
	ATOM	4473	ОН2 НОН А 926	76.929	19.455	31.644	1.00 86.93
15	ATOM	4476	ОН2 НОН А 927	171.524	31.723	13.880	1.00 59.94
	ATOM	4479	ОН2 НОН А 928	108.006	35.100	24.514	1.00 49.31
20	ATOM	4482	ОН2 НОН А 929	141.049	21.649	5.587	1.00 42.12
	MOTA	4485	он2 нон А 930	110.883	40.970	11.363	1.00 35.61
	ATOM	4488	он2 нон А 931	74.360	34.525	47.158	1.00 75.14
25	ATOM	4491	он2 нон А 932	68.751	40.894	44.809	1.00 43.22
	ATOM	4494	он2 нон А 933	59.758	29.313	23.460	1.00 61.69
30	ATOM	4497	ОН2 НОН А 934	95.173	16.341	20.293	1.00 48.05
	MOTA	4500	ОН2 НОН А 935	111.623	41.720	8.640	1.00 53.02
0.5	ATOM	4503	ОН2 НОН А 936	105.604	16.201	11.554	1.00 71.57
35	MOTA	4506	ОН2 НОН А 937	97.160	30.542	3.465	1.00 58.62
	MOTA	4509	ОН2 НОН А 938	108.492	10.703	6.225	1.00 76.16
40	ATOM	4512	ОН2 НОН А 939	135.408	44.678	13.765	1.00 61.58
	ATOM	4515	ОН2 НОН А 940	91.469	31.723	41.685	1.00 41.59
45	ATOM	4518	OH2 HOH A 941	164.580	32.799	18.975	1.00 48.97
43	ATOM	4521	ОН2 НОН А 942	157.888	29.985	20.509	1.00 63.61
	ATOM	4524	он2 нон А 943	121.776	21.112	25.392	1.00 61.93
50	ATOM	4527	он2 нон А 944	96.503	38.299	34.009	1.00 64.71
	MOTA	4530	он2 нон А 945	96.403	17.036	28.115	1.00 76.51
55	ATOM	4533	он2 нон А 946	88.114	18.204	31.407	1.00 52.33
))	MOTA	4536	он2 нон А 947	81.217	23.098	12.608	1.00 35.93
	ATOM	4539	он2 нон А 948	126.713	24.917	0.514	1.00 82.50
60	TER END						
			Atom Type Residue #	X	<u>¥</u>	<u>z</u>	OCC B

```
REMARK
     REMARK
              3 REFINEMENT.
                  PROGRAM : X-FIG.
BRUNGER
                              : X-PLOR(online) 3.843
     REMARK
     REMARK
 5
     REMARK
     REMARK
              3 DATA USED IN REFINEMENT.
                  RESOLUTION RANGE HIGH (ANGSTROMS) : 2.40
     REMARK
               3
                  RESOLUTION RANGE LOW (ANGSTROMS) :50.00
     REMARK
     REMARK
                  DATA CUTOFF
                                         (SIGMA(F)): 0.0
10
                  DATA CUTOFF HIGH
                                          (ABS(F)): 100000.00
     REMARK
                                            (ABS(F)) :
                  DATA CUTOFF LOW
     REMARK
                                                         0.010000
     REMARK
                  COMPLETENESS (WORKING+TEST ) (%): 92.7
                  NUMBER OF REFLECTIONS
     REMARK
                                                     : 18908
     REMARK
15
     REMARK
             3 FIT TO DATA USED IN REFINEMENT.
     REMARK
                  CROSS-VALIDATION METHOD
                                                   : THROUGHOUT
                  FREE R VALUE TEST SET SELECTION : RANDOM
     REMARK
                                 (WORKING SET) : 0.225
      REMARK
              3
                  R VALUE
     REMARK
                  FREE R VALUE
                                                    : 0.295
20
                  FREE R VALUE TEST SET SIZE
                                                (%): 10.2
     REMARK
                  FREE R VALUE TEST SET COUNT
     REMARK
                                                   : 1926
                  ESTIMATED ERROR OF FREE R VALUE : 0.007
      REMARK
      REMARK
     REMARK
                 FIT IN THE HIGHEST RESOLUTION BIN.
25
                  TOTAL NUMBER OF BINS USED
     REMARK
                                                           6
                  BIN RESOLUTION RANGE HIGH
                                                   (A) : 2.40
     REMARK
      REMARK
                  BIN RESOLUTION RANGE LOW
                                                   (A): 2.55
                  BIN COMPLETENESS (WORKING+TEST) (%): 94.1
      REMARK
              3
      REMARK
                  REFLECTIONS IN BIN (WORKING SET) : 2875
                                         (WORKING SET) : 0.368
                  BIN R VALUE
30
      REMARK
                  BIN FREE R VALUE
      REMARK
                                                       : 0.455
                  BIN FREE R VALUE TEST SET SIZE (%): 8.7
      REMARK
                  BIN FREE R VALUE TEST SET COUNT
      REMARK
                                                           275
      REMARK
                  ESTIMATED ERROR OF BIN FREE R VALUE : 0.027
35
      REMARK
      REMARK
               3 NUMBER OF NON-HYDROGEN ATOMS USED IN REFINEMENT.
      REMARK
                  PROTEIN ATOMS
                                           : 3531
                  NUCLEIC ACID ATOMS
      REMARK
                                                 0
                  HETEROGEN ATOMS
      REMARK
                                               102
40
      REMARK
                  SOLVENT ATOMS
                                                48
      REMARK
      REMARK
                 B VALUES.
                  FROM WILSON PLOT
                                             (A**2) : 45.0
      REMARK
                  MEAN B VALUE (OVERALL, A**2): 45.0
      REMARK
45
                   OVERALL ANISOTROPIC B VALUE.
      REMARK
                   B11 (A**2) :-11.35
      REMARK
                   B22 (A**2) : 6.33
      REMARK
               3
                   B33 (A**2) : 5.02
B12 (A**2) : 0.00
B13 (A**2) : 7.70
      REMARK
               3
      REMARK
               3
50
      REMARK
               3
                    B23 (A**2) : 0.00
      REMARK
      REMARK
               3
                  ESTIMATED COORDINATE ERROR.
      REMARK
               3
                   ESD FROM LUZZATI PLOT
                                                 (A) : 0.34
      REMARK
               3
55
                   ESD FROM SIGMAA
                                                (A) : 0.47
      REMARK
                                                (A) : 5.00
      REMARK
                   LOW RESOLUTION CUTOFF
      REMARK
               3
                  CROSS-VALIDATED ESTIMATED COORDINATE ERROR.
      REMARK
               3
                   ESD FROM C-V LUZZATI PLOT
      REMARK
                                              (A) : 0.45
60
      REMARK
                   ESD FROM C-V SIGMAA
                                                 (A) : 0.46
               3
      REMARK
               3
                  RMS DEVIATIONS FROM IDEAL VALUES.
      REMARK
                   BOND LENGTHS
      REMARK
                                                (A) : 0.007
               3
                                          (DEGREES) : 1.4
(DEGREES) : 26.1
                   BOND ANGLES
      REMARK
               3
65
                   DIHEDRAL ANGLES
      REMARK
               3
                   IMPROPER ANGLES
                                          (DEGREES) : 1.24
      REMARK
               3
      REMARK
```

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3 ISOTROPIC THERMAL MODEL : RESTRAINED
     REMARK
     REMARK
              3
                ISOTROPIC THERMAL FACTOR RESTRAINTS.
                                                        RMS
                                                                SIGMA
     REMARK
             3
                                              (A**2) : 1.77 ;
                 MAIN-CHAIN BOND
                                                               1.50
     REMARK
                 MAIN-CHAIN ANGLE
                                              (A**2) : 3.00 ; 2.00
 5
     REMARK
              3
                 SIDE-CHAIN BOND
                                              (A**2) : 2.81 ; 2.00
     REMARK
              3
                                              (A**2): 4.53; 2.50
                 SIDE-CHAIN ANGLE
     REMARK
              3
     REMARK
              3 NCS MODEL
                                       : NONE
     REMARK
10
     REMARK
             3
     REMARK
             3 NCS RESTRAINTS.
                                                       RMS
                                                           SIGMA/WEIGHT
                 GROUP 1 POSITIONAL (A): NULL; NULL
GROUP 1 B-FACTOR (A**2): NULL; NULL
     REMARK
     REMARK
     REMARK
15
              3 PARAMETER FILE 1 : parhcsdx.pro
     REMARK
              3 PARAMETER FILE 2 : param11.wat
     REMARK
                TOPOLOGY FILE 1 : tophcsdx.pro
TOPOLOGY FILE 2 : toph11.wat
     REMARK
             3
     REMARK
             3
     REMARK
              3
              3 OTHER REFINEMENT REMARKS: BULK SOLVENT MODEL USED
20
     REMARK
             1 A 507 VAL ASN PRO GLY VAL VAL ARG ILE SER GLN LYS GLY
     SEORES
             2 A 507
                       LEU ASP TYR ALA SER GLN GLN GLY THR ALA ALA LEU GLN
     SEORES
                       LYS GLU LEU LYS ARG ILE LYS ILE PRO ASP TYR SER ASP
             3 A 507
     SEQRES
             4 A
                       SER PHE LYS ILE LYS HIS LEU GLY LYS GLY HIS TYR SER
     SEQRES
                  507
25
     SEORES
             5 A
                  507
                       PHE TYR SER MET ASP ILE ARG GLU PHE GLN LEU PRO SER
                       SER GLN ILE SER MET VAL PRO ASN VAL GLY LEU LYS PHE
              6 A
                  507
     SEORES
                       SER ILE SER ASN ALA ASN ILE LYS ILE SER GLY LYS TRP
     SEQRES
             7 A
                  507
                       LYS ALA GLN LYS ARG PHE LEU LYS MET SER GLY ASN PHE
     SEORES
              8 A
                  507
             9 A
                  507
                       ASP LEU SER ILE GLU GLY MET SER ILE SER ALA ASP LEU
     SEORES
30
            10 A
                  507
                       LYS LEU GLY SER ASN PRO THR SER GLY LYS PRO THR ILE
     SEQRES
                       THR CYS SER SER CYS SER SER HIS ILE ASN SER VAL HIS
     SEQRES
             11 A
                  507
           12 A
                  507
                       VAL HIS ILE SER ALA ALA SER VAL GLY TRP LEU ILE GLN
     SECRES
                       LEU PHE HIS LYS LYS ILE GLU SER ALA LEU ARG ASN LYS
            13 A
                  507
     SEORES
                       MET ASN SER GLN VAL CYS GLU LYS VAL THR ASN SER VAL
                  507
     SEQRES
            14 A
            15 A
35
                       SER SER GLU LEU GLN PRO TYR PHE GLN THR LEU PRO VAL
     SEQRES
                  507
                       MET THR LYS ILE ASP SER VAL ALA GLY ILE ASN TYR GLY
     SEORES
             16 A
                  507
                  507
                       LEU VAL ALA PRO PRO ALA THR THR ALA GLU THR LEU ASP
            17 A
     SEQRES
     SEQRES
            18 A
                   507
                       VAL GLN MET LYS GLY GLU PHE TYR SER GLU ALA ALA ALA
                  507
                       ALA PRO PRO PRO PHE ALA PRO PRO VAL MET GLU PHE PRO
            19 A
     SEORES
           20 A
40
     SEQRES
                  507
                       ALA ALA ALA ASP ARG MET VAL TYR LEU GLY LEU SER ASP
                       TYR PHE PHE ASN THR ALA GLY LEU VAL TYR GLN GLU ALA
     SEORES
             21 A
                   507
            22 A
                   507
                       GLY VAL LEU LYS MET THR LEU ARG ASP ASP MET ILE PRO
     SEQRES
     SEORES
            23 A
                   507
                       LYS GLU SER ALA PHE ARG LEU THR THR SER PHE PHE GLY
                       THR PHE LEU PRO GLU VAL ALA LYS LYS PHE PRO ASN MET
                  507
     SEQRES
            24 A
            25 A
45
                  507
                       LYS ILE GLN ILE HIS VAL SER ALA SER THR PRO PRO HIS
     SEQRES
                       LEU SER VAL GLN PRO THR GLY LEU THR PHE TYR PRO ALA
                   507
     SEORES
            26 A
            27 A
                       VAL ASP VAL GLN ALA PHE ALA VAL LEU PRO ASN SER ALA
     SEQRES
                   507
                       LEU ALA SER LEU PHE LEU ILE GLY MET HIS THR THR GLY
     SEORES 28 A
                  507
     SEQRES 29 A
                   507
                       SER MET GLU VAL SER ALA GLU SER ASN ARG LEU VAL GLY
                       GLU LEU LYS LEU ASP ARG LEU LEU LEU GLU LEU LYS HIS
50
     SEQRES
                   507
            30 A
                       SER ASN ILE GLY PRO PHE PRO VAL GLU LEU LEU GLN ASP
     SEQRES
            31 A 507
                       ILE MET ASN TYR ILE VAL PRO ILE LEU VAL LEU PRO ARG
     SEORES
             32 A
                   507
                       VAL ASN GLU LYS LEU GLN LYS GLY PHE PRO LEU PRO THR
                   507
     SEQRES
             33 A
                       PRO ALA ARG VAL GLN LEU TYR ASN VAL VAL LEU GLN PRO
     SEQRES
             34 A
                  507
55
             35 A
                   507
                       HIS GLN ASN PHE LEU LEU PHE GLY ALA ASP VAL VAL TYR
     SEORES
                  507
                       LYS PC PC HOH HOH HOH HOH HOH HOH HOH HOH HOH
     SEQRES
             36 A
                  507
                       37 A
     SEQRES
             38 A
                       507
     SEORES
             SEQRES
             1 CYS A 135 CYS A 175
60
     SSBOND
             185.600
                       33.000 85.200 90.00 101.60 90.00 C 2
     CRYST1
                 1.000000 0.000000 0.000000
0.000000 1.000000 0.000000
0.000000 0.000000 1.000000
                                                    0.00000
     ORIGX1
     ORIGX2
                                                     0.00000
     ORIGX3
                                                    0.00000
               0.005388 0.000000 0.001106
0.000000 0.030303 0.000000
0.000000 0.000000 0.011982
65
                                                    0.00000
     SCALE1
                                                    0.00000
     SCALE2
                                                   0.00000
     SCALE3
```

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The following abbreviations are used in Table 4 in accordance with the format and usage established by the Protein Data Bank ("PDB"), Brookhaven National Laboratory, Brookhaven, N.Y. These coordinates are on deposit with the PDB, ID code 1bp1. Atomic coordinates of a BPI protein as described herein appear at Table 4 (pages 62-171 herein) and refinement statistics also appear at the end of Table 4 (pages 172-173 herein). Table 4 corresponds to Figure 6 (Figs. 6.1-6.112) in United States patent application serial no. 08/879,565, filed June 20, 1997.

"Atom type" refers to the element whose coordinates are measured. The first letter in the column defines the element.

"Residue" refers to the amino acid in the BPI protein sequence, using the standard three letter abbreviations known in the art.

"#" refers to the residue number.

"X, Y, Z" crystallographically define the atomic position, in three-dimensional space, of the element measured.

"OCC" is the occupancy value.

"B" is a thermal factor that measures movement of the atom around its atomic center.

10. Organomimetics

Molecular modelling of BPI as described herein is useful for the preparation of organomimetics such as "surface" mimetics. As one example, organomimetics are prepared based on "tip" mimetics in which the three-dimensional coordinates of the tip, as described above, are used to create a "surface" (or complementary pocket) into which a computer program builds an organic molecule with similar characteristics.

Numerous modifications and variations in the practice of the invention are expected to occur to those skilled in the art upon consideration of the foregoing description and the presently preferred embodiments thereof. Consequently, the scope of the present invention is to be defined by the appended claims.

SEQUENCE LISTING

- (1) GENERAL INFORMATION:
- (i) APPLICANT: XOMA Corporation; (Beamer, Lesa J., Carroll, Stephen F., Eisenberg, David, U.S. only)
- (ii) TITLE OF INVENTION: Bactericidal/Permeability-Increasing Protein: Crystallization, X-Ray Diffraction, Three-Dimensional Structure Determination, Rational Drug Design and Molecular Modeling of Related Proteins
 - (iii) NUMBER OF SEQUENCES: 2
 - (iv) CORRESPONDENCE ADDRESS:
 - (A) ADDRESSEE: McAndrews, Held & Malloy
 - (B) STREET: 500 West Madison, 34th Floor

 - (C) CITY: Chicago (D) STATE: Illinois
 - (E) COUNTRY: United States of America
 - (F) ZIP: 60661
 - (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.25
 - (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER:
 - (B) FILING DATE:
 - (C) CLASSIFICATION:
 - (vii) PRIOR APPLICATION DATA:
 - (A) APPLICATION NUMBER: U.S. Serial No. 08/879,565
 (B) FILING DATE: June 20, 1997

 - (C) CLASSIFICATION:
 - (viii) ATTORNEY/AGENT INFORMATION:
 - (A) NAME: McNicholas, Janet M.
 - (B) REGISTRATION NUMBER: 32,918
 - (C) REFERENCE/DOCKET NUMBER: 11034W001/100-248.PCT
 - (ix) TELECOMMUNICATION INFORMATION:
 - (A) TELEPHONE: 312/707-8889
 - (B) TELEFAX: 312/707-9155
 - (C) TELEX:
- (2) INFORMATION FOR SEQ ID NO:1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1813 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: cDNA
 - (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION: 31..1491
 - (ix) FEATURE:
 - (A) NAME/KEY: mat_peptide

(B) LOCATION: 124..1491

- (ix) FEATURE:
 (A) NAME/KEY: misc_feature
 - (D) OTHER INFORMATION: "rBPI"

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

	(XI) SEQUENCE DESCRIPTION: SEQ ID NO:1:															
CAGG	CCTI	'GA G	GTTI	TGGC	A GC	TCTG	GAGG	Met		g Glu			-		GGC	54
			GCG Ala -20													102
			GTG Val													150
			GGC Gly													198
			CTG Leu													246
			CAT His 45													294
			TTC Phe													342
			AAG Lys													390
			GCA Ala													438
			GAA Glu							Asp						486
			TCA Ser 125													534
			AGT Ser					Ile					Val			582
CTG Leu	ATC Ile 155	CAA Gln	CTC Leu	TTC Phe	CAC His	AAA Lys 160	Lys	ATT	GAG Glu	TCT Ser	GCG Ala 165	Leu	CGA Arg	AAC Asn	AAG Lys	630
			CAG Gln			Glu					Ser					678
CTG	CAA	CCT	TAT	TTC	CAG	ACT	CTG	CCA	GTA	ATG	ACC	AAA	ATA	GAT	TCT	726

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Leu	Gln	Pro	Tyr	Phe 190	Gln	Thr	Leu	Pro	Val 195	Met	Thr	Lys	Ile	Asp 200	Ser	
GTG Val	GCT Ala	GGA Gly	ATC Ile 205	AAC Asn	TAT Tyr	GGT Gly	CTG Leu	GTG Val 210	GCA Ala	CCT Pro	CCA Pro	GCA Ala	ACC Thr 215	ACG Thr	GCT Ala	774
GAG Glu	ACC Thr	CTG Leu 220	GAT Asp	GTA Val	CAG Gln	ATG Met	AAG Lys 225	GGG Gly	GAG Glu	TTT Phe	TAC Tyr	AGT Ser 230	GAG Glu	AAC Asn	CAC His	822
CAC His	AAT Asn 235	CCA Pro	CCT Pro	CCC Pro	TTT Phe	GCT Ala 240	CCA Pro	CCA Pro	GTG Val	ATG Met	GAG Glu 245	TTT Phe	CCC Pro	GCT Ala	GCC Ala	870
CAT His 250	GAC Asp	CGC Arg	ATG Met	GTA Val	TAC Tyr 255	CTG Leu	GGC Gly	CTC Leu	TCA Ser	GAC Asp 260	TAC Tyr	TTC Phe	TTC Phe	AAC Asn	ACA Thr 265	918
GCC Ala	GGG Gly	CTT Leu	GTA Val	TAC Tyr 270	CAA Gln	GAG Glu	GCT Ala	GGG Gly	GTC Val 275	TTG Leu	AAG Lys	ATG Met	ACC Thr	CTT Leu 280	AGA Arg	966
GAT Asp	GAC Asp	ATG Met	ATT Ile 285	CCA Pro	AAG Lys	GAG Glu	TCC Ser	AAA Lys 290	TTT Phe	CGA Arg	CTG Leu	ACA Thr	ACC Thr 295	AAG Lys	TTC Phe	1014
TTT Phe	GGA Gly	ACC Thr 300	TTC Phe	CTA Leu	CCT Pro	GAG Glu	GTG Val 305	GCC Ala	AAG Lys	AAG Lys	TTT Phe	CCC Pro 310	AAC Asn	ATG Met	AAG Lys	1062
ATA Ile	CAG Gln 315	ATC Ile	CAT His	GTC Val	TCA Ser	GCC Ala 320	TCC Ser	ACC Thr	CCG Pro	CCA Pro	CAC His 325	CTG Leu	TCT Ser	GTG Val	CAG Gln	1110
CCC Pro 330	ACC Thr	GGC Gly	CTT Leu	ACC Thr	TTC Phe 335	TAC Tyr	CCT Pro	GCC Ala	GTG Val	GAT Asp 340	GTC Val	CAG Gln	GCC Ala	TTT Phe	GCC Ala 345	1158
GTC Val	CTC Leu	CCC Pro	AAC Asn	TCC Ser 350	TCC Ser	CTG Leu	GCT Ala	TCC Ser	CTC Leu 355	TTC Phe	CTG Leu	ATT Ile	GGC Gly	ATG Met 360	CAC His	1206
ACA Thr	ACT Thr	GGT Gly	TCC Ser 365	ATG Met	GAG Glu	GTC Val	AGC Ser	GCC Ala 370	GAG Glu	TCC Ser	AAC Asn	AGG Arg	CTT Leu 375	GTT Val	GGA Gly	1254
GAG Glu	CTC Leu	AAG Lys 380	CTG Leu	GAT Asp	AGG Arg	CTG Leu	CTC Leu 385	CTG Leu	GAA Glu	CTG Leu	AAG Lys	CAC His 390	TCA Ser	AAT Asn	ATT Ile	1302
GGC Gly	CCC Pro 395	TTC Phe	CCG Pro	GTT Val	GAA Glu	TTG Leu 400	CTG Leu	CAG Gln	GAT Asp	ATC Ile	ATG Met 405	AAC Asn	TAC Tyr	ATT Ile	GTA Val	1350
CCC Pro 410	Ile	CTT	GTG Val	CTG Leu	CCC Pro 415	AGG Arg	GTT Val	AAC Asn	GAG Glu	AAA Lys 420	CTA Leu	CAG Gln	AAA Lys	GC	TTC Phe 425	1398
CCT Pro	CTC Leu	CCG Pro	ACG Thr	CCG Pro 430	GCC Ala	AGA Arg	GTC Val	CAG Gln	CTC Leu 435	TAC Tyr	AAC Asn	GTA Val	GTG Val	CTT Leu 440	CAG Gln	1446
CCT Pro	CAC His	CAG Gln	AAC Asn 445	TTC Phe	CTG Leu	CTG Leu	TTC Phe	GGT Gly 450	GCA Ala	GAC Asp	GTT Val	GTC Val	TAT Tyr 455	AAA Lys		1491

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TGAAGGCACC	AGGGGTGCCG	GGGGCTGTCA	GCCGCACCTG	TTCCTGATGG	GCTGTGGGGC	1551
ACCGGCTGCC	TTTCCCCAGG	GAATCCTCTC	CAGATCTTAA	CCAAGAGCCC	CTTGCAAACT	1611
TCTTCGACTC	AGATTCAGAA	ATGATCTAAA	CACGAGGAAA	CATTATTCAT	TGGAAAAGTG	1671
CATGGTGTGT	ATTTTAGGGA	TTATGAGCTT	CTTTCAAGGG	CTAAGGCTGC	AGAGATATTT	1731
CCTCCAGGAA	TCGTGTTTCA	ATTGTAACCA	AGAAATTTCC	ATTTGTGCTT	CATGAAAAA	1791
AACTTCTGGT	TTTTTTCATG	TG				1813

(2) INFORMATION FOR SEQ ID NO:2:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 487 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2: Met Arg Glu Asn Met Ala Arg Gly Pro Cys Asn Ala Pro Arg Trp Val -31 -30 Ser Leu Met Val Leu Val Ala Ile Gly Thr Ala Val Thr Ala Ala Val Asn Pro Gly Val Val Val Arg Ile Ser Gln Lys Gly Leu Asp Tyr Ala Ser Gln Gln Gly Thr Ala Ala Leu Gln Lys Glu Leu Lys Arg Ile Lys Ile Pro Asp Tyr Ser Asp Ser Phe Lys Ile Lys His Leu Gly Lys Gly . 35 40 45 His Tyr Ser Phe Tyr Ser Met Asp Ile Arg Glu Phe Gln Leu Pro Ser Ser Gln Ile Ser Met Val Pro Asn Val Gly Leu Lys Phe Ser Ile Ser 70 75 80 Asn Ala Asn Ile Lys Ile Ser Gly Lys Trp Lys Ala Gln Lys Arg Phe Leu Lys Met Ser Gly Asn Phe Asp Leu Ser Ile Glu Gly Met Ser Ile 105 Ser Ala Asp Leu Lys Leu Gly Ser Asn Pro Thr Ser Gly Lys Pro Thr Ile Thr Cys Ser Ser Cys Ser Ser His Ile Asn Ser Val His Val His 135 Ile Ser Lys Ser Lys Val Gly Trp Leu Ile Gln Leu Phe His Lys Lys

Ile Glu Ser Ala Leu Arg Asn Lys Met Asn Ser Gln Val Cys Glu Lys

Val Thr Asn Ser Val Ser Ser Lys Leu Gln Pro Tyr Phe Gln Thr Leu

Pro Val Met Thr Lys Ile Asp Ser Val Ala Gly Ile Asn Tyr Gly Leu

205 200 195 Val Ala Pro Pro Ala Thr Thr Ala Glu Thr Leu Asp Val Gln Met Lys 215 220 Gly Glu Phe Tyr Ser Glu Asn His His Asn Pro Pro Pro Phe Ala Pro Pro Val Met Glu Phe Pro Ala Ala His Asp Arg Met Val Tyr Leu Gly Leu Ser Asp Tyr Phe Phe Asn Thr Ala Gly Leu Val Tyr Gln Glu Ala Gly Val Leu Lys Met Thr Leu Arg Asp Asp Met Ile Pro Lys Glu Ser Lys Phe Arg Leu Thr Thr Lys Phe Phe Gly Thr Phe Leu Pro Glu Val 295 Ala Lys Lys Phe Pro Asn Met Lys Ile Gln Ile His Val Ser Ala Ser Thr Pro Pro His Leu Ser Val Gln Pro Thr Gly Leu Thr Phe Tyr Pro Ala Val Asp Val Gln Ala Phe Ala Val Leu Pro Asn Ser Ser Leu Ala Ser Leu Phe Leu Ile Gly Met His Thr Thr Gly Ser Met Glu Val Ser Ala Glu Ser Asn Arg Leu Val Gly Glu Leu Lys Leu Asp Arg Leu Leu Leu Glu Leu Lys His Ser Asn Ile Gly Pro Phe Pro Val Glu Leu Leu 390 Gln Asp He Met Asn Tyr Ile Val Pro Ile Leu Val Leu Pro Arg Val Asn Glu Lys Leu Gln Lys Gly Phe Pro Leu Pro Thr Pro Ala Arg Val Gln Leu Tyr Asn Val Val Leu Gln Pro His Gln Asn Phe Leu Leu Phe Gly Ala Asp Val Val Tyr Lys

CLAIMS

What Is Claimed Is:

- 1. Use of atomic coordinates of bactericidal/permeability-increasing ("BPI") protein, or fragment, analog or variant thereof, to model a BPI protein.
- 2. Use of atomic coordinates of bactericidal/permeability-increasing ("BPI") protein, or fragment, analog or variant thereof, to model a BPI-related lipid transfer protein.
- 3. The use according to claim 2, wherein the BPI-related lipid transfer protein is lipopolysaccharide-binding protein (LBP), cholesteryl ester transferase protein (CETP) or phospholipid transfer protein (PLTP), or fragment, analog or variant thereof.
- 4. The use according to any of claims 1-3, wherein the BPI protein comprises a binding site characterized by amino acid residues of at least one binding pocket as defined in Table 31.
- 5. The use according to any of claims 1-3, wherein the BPI protein comprises a binding site characterized by at least one amino acid sequence, or variant of the sequence, selected from positions about 17 to about 45, positions about 36 to about 54, positions about 65 to about 99, positions about 84 to about 109, positions about 142 to about 164, or positions about 142 to about 169 of BPI.
- 6. The use according to any of claims 1-3, wherein the BPI protein comprises a binding site characterized by amino acid residues of at least one binding pocket as defined in Table 3 and a binding site characterized by at least one amino acid sequence, or variant of the sequence, selected from positions about 17 to about 45, positions about 36 to about 54, positions about 65 to about 99, positions about 84 to about 109, positions about 142 to about 164, or positions about 142 to about 169 of BPI.

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- 7. Use of atomic coordinates of bactericidal/permeability-increasing ("BPI") protein to computationally design a chemical compound for mimicking BPI protein, or fragment, analog or variant thereof.
- 8. Use of atomic coordinates of bactericidal/permeability-increasing ("BPI") protein to computationally design a chemical compound for mimicking a BPI-related lipid transfer protein, or fragment, analog or variant thereof.
- 9. The use according to claim 8, wherein the BPI-related lipid transfer protein is lipopolysaccharide-binding protein (LBP), cholesteryl ester transferase protein (CETP) or phospholipid transfer protein (PLTP).
- 10. Use of atomic coordinates of bactericidal/permeability-increasing ("BPI") protein, to design a chemical compound capable of associating with a BPJ-related lipid binding protein, or fragment, analog or variant thereof.
- 11. The use according to claim 10, wherein the BPI-related lipid binding protein is lipopolysaccharide-binding protein (LBP), cholesteryl ester transferase protein (CETP) or phospholipid transfer protein (PLTP), or fragment, analog or variant thereof.
- 12. Use of atomic coordinates of bactericidal/permeability-increasing ("BPI") protein to design a model of ligands in an active site of a lipid binding protein.
- 13. The use according to claim 12, wherein the lipid binding protein is bactericidal/permeability-increasing protein (BPI), lipopolysaccharide-binding protein (LBP), cholesteryl ester transferase protein (CETP) or phospholipid transfer protein (PLTP), or fragment, analog or variant thereof.
- 14: Use of atomic coordinates of bactericidal/permeability-increasing ("BPI") protein, to design compounds with at least one activity selected from the group consisting of antibacterial, antifungal, antimycobacterial, antichlamydial, antiprotozoan, heparin-binding, endotoxin-binding, heparin-neutralizing, endotoxin-neutralizing, inhibition of tumor and

endothelial cell proliferation, inhibition of angiogenesis, anti-inflammatory, anticoagulant and antithrombolytic.

- 15. The use according to any of claims 1 14, wherein said atomic coordinates are according to Table 4.
- 16. A method of three-dimensional modeling of a bactericidal/permeability-increasing ("BPI") protein comprising the steps of:
 - (a) providing three-dimensional atomic coordinates derived from X-ray diffraction measurements of a BPI protein in a computer readable format;
 - (b) inputting the data from step (a) into a computer with appropriate software programs;
 - (c) generating a three-dimensional structural representation of the BPI protein suitable for visualization and further computational manipulation.
- 17. A method of three-dimensional modeling of a bactericidal/permeability-increasing ("BPI")-related lipid transfer protein comprising the steps of:
 - (a) providing three-dimensional atomic coordinates derived from X-ray diffraction measurements of a BPI protein in a computer readable format;
 - (b) inputting the data from step (a) into a computer with appropriate software programs;
 - (c) generating a three-dimensional structural representation of the BPI-related lipid transfer protein suitable for visualization and further computational manipulation.
- 18. The use according to any of claims 16-17, wherein the BPI protein comprises a binding site characterized by amino acid residues of at least one binding pocket as defined in Table 3.
- 19. The use according to any of claims 16-17, wherein the BPI protein comprises a binding site characterized by at least one amino acid sequence, or variant of the sequence, selected from positions about 17 to about 45, positions about 36 to about 54, positions about

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65 to about 99, positions about 84 to about 109, positions about 142 to about 164, or positions about 142 to about 169 of BPI.

- 20. The use according to any of claims 16-17, wherein the BPI protein comprises a binding site characterized by amino acid residues of at least one binding pocket as defined in Table 3 and a binding site characterized by at least one amino acid sequence, or variant of the sequence, selected from positions about 17 to about 45, positions about 36 to about 54, positions about 65 to about 99, positions about 84 to about 109, positions about 142 to about 164, or positions about 142 to about 169 of BPI.
- 21. A method for providing an atomic model of a BPI protein, or fragment, analog or variant thereof, comprising
 - (a) providing a computer readable medium having stored thereon atomic coordinate/x-ray diffraction data of the BPI protein, or fragment, analog or variant thereof, in crystalline form, the data sufficient to model the three-dimensional structure of the BPI protein, or fragment, analog or variant thereof;
 - (b) analyzing, on a computer using at least one subroutine executed in said computer, atomic coordinate/x-ray diffraction data from (a) to provide atomic coordinate data output defining an atomic model of said BPI protein, or fragment, analog or variant thereof, said analyzing utilizing at least one computing algorithm selected from the group consisting of data processing and reduction, auto-indexing, intensity scaling, intensity merging, amplitude conversion, truncation, molecular replacement, molecular alignment, molecular refinement, electron density map calculation, electron density modification, electron map visualization, model building, rigid body refinement, positional refinement; and
 - (c) obtaining atomic coordinate data defining the three-dimensional structure of at least one of said BPI protein, or fragment, analog or variant thereof.

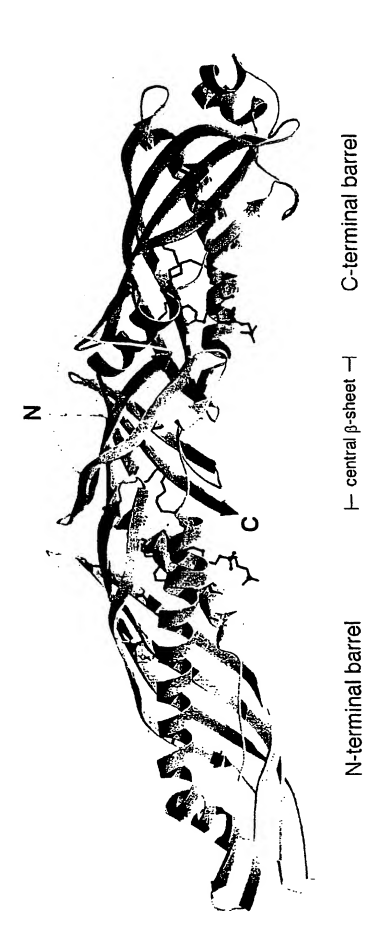
- 22. A method according to claim 21, wherein said computer readable medium further has stored thereon data corresponding to a nucleic acid sequence or an amino acid sequence data comprising at least one structural domain or functional domain of a BPI, LBP, CETP or PLTP corresponding to at least one BPI or mutant primary sequence of Figures 2-20 or Table 2, or a fragment thereof; and wherein said analyzing step further comprises analyzing said sequence data.
- 23. A computer-based system for providing atomic model data of the three-dimensional structure of BPI protein, or fragment, analog or variant thereof, a BPI mutant or a BPI fragment, comprising the following elements:
 - at least one computer readable medium (CRM) having stored thereon atomic coordinate/x-ray diffraction data of said BPI protein, or fragment, analog or variant thereof;
 - (b) at least one computing subroutine that, when executed in a computer, causes the computer to analyze atomic coordinate/x-ray diffraction data from (a) to provide atomic coordinate data output defining an atomic model of said BPI protein, or fragment, analog or variant thereof, said analyzing utilizing at least one computing subroutine selected from the group consisting of data processing and reduction, auto-indexing, intensity scaling, intensity merging, amplitude conversion, truncation, molecular replacement, molecular alignment, molecular refinement, electron density map calculation, electron density modification, electron map visualization, model building, rigid body refinement, positional refinement; and
 - (c) retrieval means for obtaining atomic coordinate output data substantially defining the three-dimensional structure of said BPI protein, or fragment, analog or variant thereof.
- 24. A method for providing a computer atomic model of a ligand of a BPI protein, or fragment, analog or variant thereof, comprising

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- (a) providing a computer readable medium (CRM) having stored thereon atomic coordinate data of a BPI protein, or fragment, analog, or variant thereof;
- (b) providing a CRM having stored thereon atomic coordinate data sufficient to generate atomic models of potential ligands of said BPI protein, or fragment, analog, or variant thereof;
- (c) analyzing on a computer, using at least one subroutine executed in said computer, the atomic coordinate data from (a) and ligand data from (b), to determine binding sites of BPI protein, or fragment, analog, or variant thereof, and to provide atomic coordinate data defining an atomic model of at least one ligand of said BPI, BPI mutant or a fragment thereof, said analyzing utilizing computing subroutines selected from the group consisting of data processing and reduction, auto-indexing, intensity scaling, intensity merging, amplitude conversion, truncation, molecular replacement, molecular alignment, molecular refinement, electron density map calculation, electron density modification, electron map visualization, model building, rigid body refinement, positional refinement; and
- (d) obtaining atomic coordinate model output data defining the threedimensional structure of said at least one ligand of said BPI protein, or fragment, analog, or variant thereof.
- 25. A computer-based system for providing an atomic model of at least one ligand of a BPI, BPI mutant or a fragment thereof, comprising the following elements;
 - (a) a computer readable medium (CRM) having stored thereon atomic coordinate data of a BPI, mutant or fragment thereof;
 - (b) a CRM having stored thereon atomic coordinate data sufficient to generate atomic models of potential ligands of a BPI, mutant or fragment;
 - (c) at least one computing subroutine for analyzing on a computer, the atomic coordinate data from (a) and (b), to determine binding sites of BPI protein, or fragment, analog, or variant thereof, and to provide

data output defining an atomic model of at least one potential ligand of BPI protein, or fragment, analog, or variant thereof, said analyzing utilizing at least one computing subroutine selected from the group consisting of data processing and reduction, auto-indexing, intensity scaling, intensity merging, amplitude conversion, truncation, molecular replacement, molecular alignment, molecular refinement, electron density map calculation, electron density modification, electron map visualization, model building, rigid body refinement, positional refinement; and

(d) retrieval means for obtaining atomic coordinate data of said at least one ligand of a BPI protein, or fragment, analog or variant thereof.



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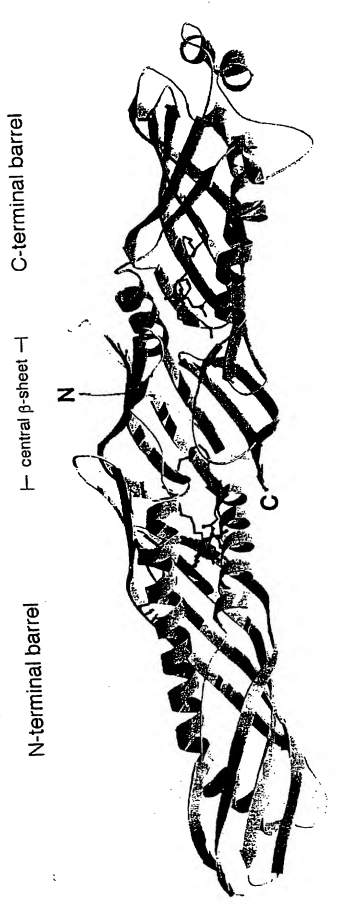


Fig. 1B

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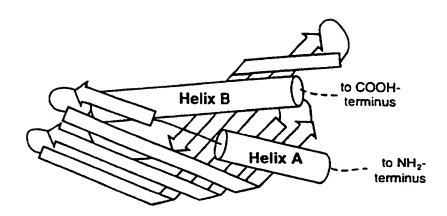
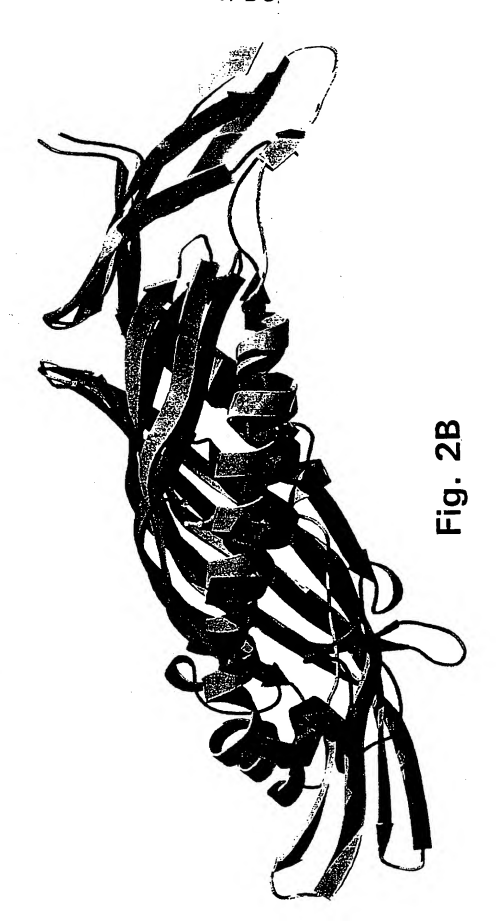
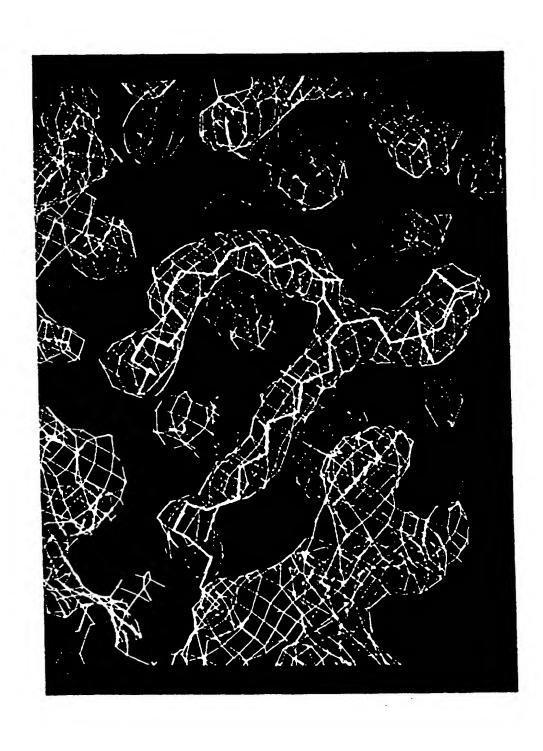
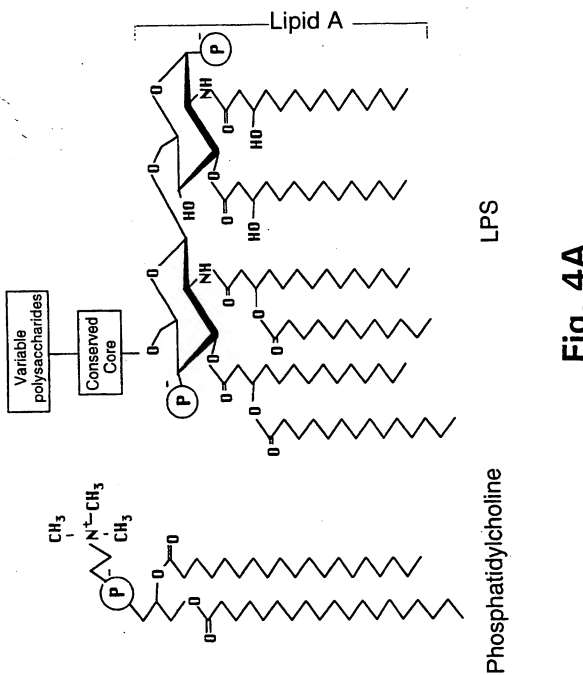


Fig. 2A

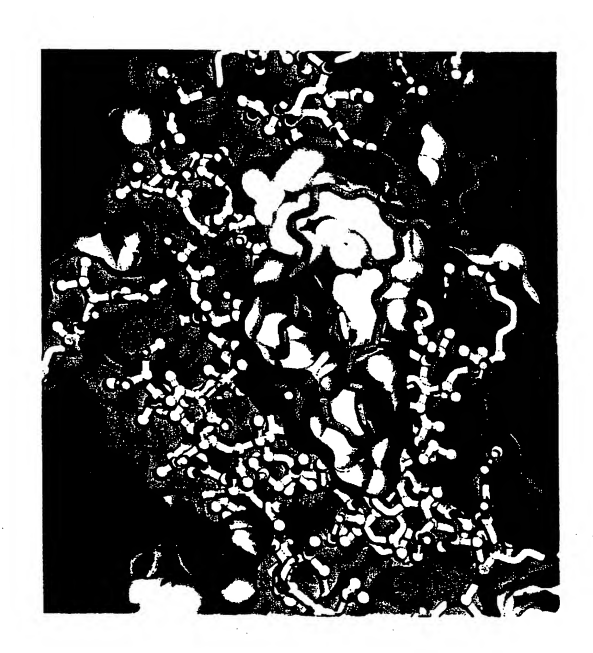








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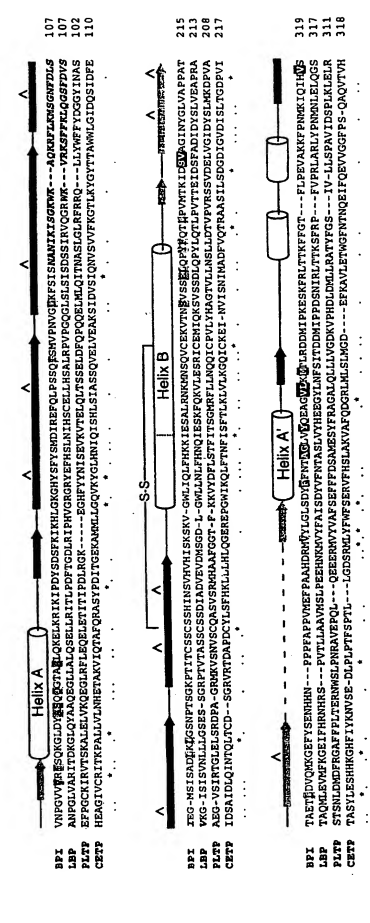


Fig. 5A

411 409 403 409	
Helix B' LODIMNYIVPILQDIMNYIVPILQDIMNYIUN	
PVEL NAEL ALIP	456 476 470
A A A A A A A A A A A A A A A A B A B A	
BPI LBP PLTP CETP	BPI LBP PLTP CRTP

Fig. 5B

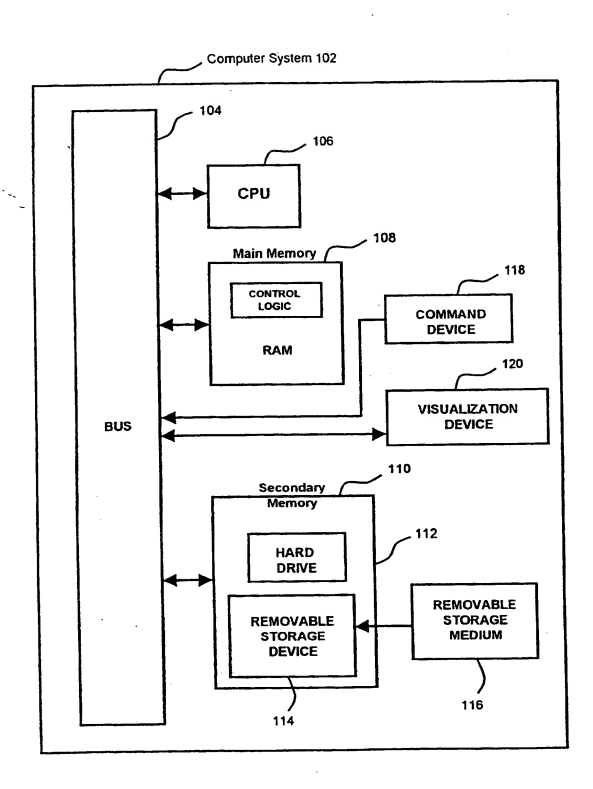


Fig. 6

INTERNATIONAL SEARCH REPORT

Int tional Application No PCT/US 98/13007

		PC1/US 98	3/ 1300/
A. CLASSII IPC 6	FICATION OF SUBJECT MATTER C07K14/47 C07K1/00 G06F17/5	0	
According to	International Patent Classification (IPC) or to both national classificat	ion and IPC	
B. FIELDS	SEARCHED		
	cumentation searched (classification system followed by classification CO7K GO1F GO6F	n symbols)	
Documentat	ion searched other than minimum documentation to the extent that su	ch documents are ancluded in the fields so	earched
Electronic da	ata base consulted during the international search (name of data base	e and, where practical, search terms used	1)
C. DOCUME	ENTS CONSIDERED TO BE RELEVANT		
Category °	Citation of document, with indication, where appropriate, of the relevant	vant passages	Relevant to claim No.
A	P W GRAY ET AL.: "Cloning of the a human neutrophil bactericidal programmed by Journal OF BIOLOGICAL CHEMISTRY., vol. 264, no. 16, 5 June 1989, pas 9505-9509, XP002034856 MD US	rotein"	1-25
Ρ,Χ	see the whole document File Medline, abstract no. 973344 XP002082313	42, 1997	1-25
	& L J BEAMER ET AL.: "Crystal st of human BPI and two bound phosph at 2.4 angstrom resolution " SCIENCE, vol. 276, no. 5320, 20 June 1997, 1861-1864, LANCASTER, PA US see abstract	olipids	
	<u> </u>	-/	
χ Furt	her documents are listed in the continuation of box C.	Patent family members are listed	d in annex.
"A" docum const "E" earlier filing "L" docum which citatic "O" docum other "P" docum	ent defining the general state of the art which is not dered to be of particular relevance document but published on or after the international date ent which may throw doubts on priority claim(s) or is cited to establish the publication date of another on or other epecial reason (as specified) ent referring to an oral disclosure, use, exhibition or means	"T" later document published after the interpretation or priority date and not in conflict with cited to understand the principle or invention. "X" document of particular relevance; the cannot be considered novel or cannot involve an inventive step when the cannot be considered to involve an document is combined with one or ments, such combination being obvin the art. "&" document member of the same pater.	th the application but theory underlying the claimed invention of be considered to document is taken alone claimed invention inventive step when the nore other such doculous to a person skilled
	actual completion of theinternational search	Date of mailing of the international s	
	27 October 1998	11/11/1998	•
Name and	mailing address of the ISA European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk	Authorized officer	

INTERNATIONAL SEARCH REPORT

Int Int Internal Application No PCT/US 98/13007

		PCT/US 98/13007					
	Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT						
Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.					
A	File Medline, abstract no. 93104264, 1993 XP002082316 & J MBONMATIN ET AL.: "Progress in multidimensional NMR investigations of peptid and protein 3-D structures in solution. From structure to functional aspects" BIOCHIMIE, vol. 74, no. 9-10, October 1992, pages 825-836, see abstract	1-25					

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